## STN-Registry/Caplus Structure Search

10/520,800

06/28/2006

```
=> s glorius f?/au
L65
           24 GLORIUS F?/AU
=> s L65 and L50
           6 L65 AND L50
=> s L65 and L63
             4 L65 AND L63
L67
=> s L66 and L67
L68
             3 L66 AND L67
=> s L66 or L50
L69
            63 L66 OR L50
=> d his
     (FILE 'HOME' ENTERED AT 09:32:44 ON 28 JUN 2006)
     FILE 'REGISTRY' ENTERED AT 09:32:52 ON 28 JUN 2006
        1650308 S NCNC2/ESS
L1
     FILE 'REGISTRY' ENTERED AT 09:51:45 ON 28 JUN 2006
L2
                STRUCTURE UPLOADED
L3
           8980 S L2 FULL
L4
                STRUCTURE UPLOADED
           3519 S L4 FULL
L5
             50 S L4
L6
L7
             50 S L2
                SAVE TEMP L5 GLOR800STR2/A
     FILE 'HCAPLUS' ENTERED AT 10:08:22 ON 28 JUN 2006
L8
            253 S L5
     FILE 'REGISTRY' ENTERED AT 10:08:52 ON 28 JUN 2006
L9
            628 S NC>1 AND L5
     FILE 'STNGUIDE' ENTERED AT 10:12:24 ON 28 JUN 2006
     FILE 'REGISTRY' ENTERED AT 10:14:37 ON 28 JUN 2006
                STRUCTURE UPLOADED
L10
L11
             50 S L10 SAM SSS SUB=L5
L12
           1426 S L10 FULL SSS SUB=L5
     FILE 'HCAPLUS' ENTERED AT 10:17:32 ON 28 JUN 2006
L13
           108 S L12
     FILE 'REGISTRY' ENTERED AT 10:17:46 ON 28 JUN 2006
L14
            237 S L12 AND NRRS>2
     FILE 'HCAPLUS' ENTERED AT 10:27:31 ON 28 JUN 2006
L15
             56 S L14
     FILE 'REGISTRY' ENTERED AT 10:27:55 ON 28 JUN 2006
L16
           1189 S L12 NOT L14
     FILE 'HCAPLUS' ENTERED AT 10:28:11 ON 28 JUN 2006
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10/520,800 06/28/2006

60 S L16 L17 8 S L15 AND L17 L18 FILE 'REGISTRY' ENTERED AT 10:29:05 ON 28 JUN 2006 FILE 'STNGUIDE' ENTERED AT 10:29:18 ON 28 JUN 2006 FILE 'REGISTRY' ENTERED AT 10:45:47 ON 28 JUN 2006 STRUCTURE UPLOADED L19 L20 32 S L19 SAM SSS SUB=L12 551 S L19 FULL SSS SUB=L12 L21 FILE 'HCAPLUS' ENTERED AT 10:52:12 ON 28 JUN 2006 L22 85 S L21 FILE 'REGISTRY' ENTERED AT 10:53:23 ON 28 JUN 2006 L23 368 S L21 NOT L14 FILE 'HCAPLUS' ENTERED AT 10:53:51 ON 28 JUN 2006 L24 45 S L23 FILE 'REGISTRY' ENTERED AT 10:54:25 ON 28 JUN 2006 L25 875 S L12 NOT L21 821 S L12 NOT (L21 OR L14) L26 3310 S 180.306.6/RID L27 809 S L26 AND L27 L28 L29 12 S L26 NOT L28 FILE 'STNGUIDE' ENTERED AT 11:02:16 ON 28 JUN 2006 FILE 'REGISTRY' ENTERED AT 11:04:38 ON 28 JUN 2006 L30 STRUCTURE UPLOADED L31 1 S L30 SAM SSS SUB=L12 35 S L30 FULL SSS SUB=L12 L32 0 S L32 AND L24 L33 35 S L32 AND L14 L34 FILE 'HCAPLUS' ENTERED AT 11:07:03 ON 28 JUN 2006 L35 11 S L34 FILE 'REGISTRY' ENTERED AT 11:07:31 ON 28 JUN 2006 FILE 'HCAPLUS' ENTERED AT 11:08:14 ON 28 JUN 2006 54 S L35 OR L24 L36 FILE 'REGISTRY' ENTERED AT 11:11:52 ON 28 JUN 2006 L37 403 S L23 OR L32 FILE 'HCAPLUS' ENTERED AT 11:13:36 ON 28 JUN 2006 1 S US2005-520800/APPS L38 SEL RN FILE 'REGISTRY' ENTERED AT 11:14:26 ON 28 JUN 2006 109 S E1-E109 L39

34 S L39 AND L37

75 S L39 NOT L40

9 S L14 AND L39

L40 L41

L42

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L43
           202 S L14 NOT L32
     FILE 'HCAPLUS' ENTERED AT 11:26:41 ON 28 JUN 2006
L44
            47 S L43
     FILE 'REGISTRY' ENTERED AT 11:27:00 ON 28 JUN 2006
     FILE 'STNGUIDE' ENTERED AT 11:31:48 ON 28 JUN 2006
    FILE 'REGISTRY' ENTERED AT 11:35:19 ON 28 JUN 2006
L45
               STRUCTURE UPLOADED
              2 S L45 SAM SSS SUB=L12
L46
             46 S L45 FULL SSS SUB=L12
L47
             46 S L14 AND L47
L48
     FILE 'HCAPLUS' ENTERED AT 11:37:27 ON 28 JUN 2006
L49
             12 S L48
             63 S L24 OR L35 OR L49
L50
     FILE 'REGISTRY' ENTERED AT 11:38:49 ON 28 JUN 2006
L51
           191 S L14 NOT L47
           161 S L14 NOT (L47 OR L32)
L52
     FILE 'REGISTRY' ENTERED AT 11:52:20 ON 28 JUN 2006
                SAVE TEMP L23 GLOR800L23/A
                SAVE TEMP L34 GLOR800L34/A
                SAVE TEMP L48 GLOR800L48/A
     FILE 'HCAPLUS' ENTERED AT 11:55:09 ON 28 JUN 2006
               SAVE TEMP L50 GLOR800L50/A
     FILE 'CASREACT' ENTERED AT 12:03:57 ON 28 JUN 2006
L53
               STRUCTURE UPLOADED
L54
             1 S L53 SAM SSS
          113 S L53 FULL SSS
L55
           85 S L55/COM
L56
               STRUCTURE UPLOADED
L57
L58
            1 S L57 SAM SSS
L59
            8 S L57 FULL SSS
L60
             STRUCTURE UPLOADED
             1 S L60
L61
             35 S L60 FULL
L62
     FILE 'HCAPLUS' ENTERED AT 12:20:00 ON 28 JUN 2006
             35 S L62
L63
             3 S L50 AND L63
L64
             24 S GLORIUS F?/AU
L65
             6 S L65 AND L50
L66
             4 S L65 AND L63
L67
             3 S L66 AND L67
L68
             63 S L66 OR L50
L69
=> file registry
                                                 SINCE FILE TOTAL ENTRY SESSION 27.83 1055.64
COST IN U.S. DOLLARS
```

FULL ESTIMATED COST

10/520,800 06/28/2006

FILE COVERS 1907 - 28 Jun 2006 VOL 145 ISS 1 FILE LAST UPDATED: 27 Jun 2006 (20060627/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d ibib abs hitstr L69 1-63

```
L69 ANSWER 1 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2006:284758 HCAPLUS DOCUMENT NUMBER: 145:7488
DOCUMENT NUMBER:
                                                                      145:7488
The first palladium-catalyzed Sonogashira coupling of unactivated secondary alkyl bromides
Altenhoff, Gereon; Wuertz, Sebastian; Glorius,
AUTHOR (S):
                                                                    Altenhoff, Gerenn; muert, statement from Frank
BASF AG, GCB/K-M311, Ludwigshafen, 67056, Germany
Tetrahedron Letters (2006), 47(17), 2925-2928
CODEN: TELEAY; ISSN: 0040-4039
Elsevier B.V.
CORPORATE SOURCE:
SOURCE:
PUBLISHER:
            LISECUTE TYPE: Journal
UAGE: English
A palladium-carbene catalyzed Sonogashira coupling of unactivated alkyl
bromides with alkyl substituted alkynes is reported. E.g.,
[(IBiox7)PdC12]2 catalyzed the Sonogashira coupling of cycloheptyl
DOCUMENT TYPE:
LANGUAGE:
             with 1-octyme to give 76% 1-octynylcycloheptane. For the first time, unactivated secondary alkyl halides were successfully employed in Sonogashira reactions. 606970-69-8
            606970-69-8
RL: CAT (Catelyst use); USES (Uses)
    (palladium-carbene catalyzed Sonogashira coupling of unactivated alkyl bromides with alkyl substituted alkynes)
606970-69-8 HCAPLUS
Dispiro(cyclohexane-1,3'(2'H)-imidazo{5,1-b:4,3-b'}bisoxazol{4}ium-7'(8'H),1''-cyclohexane}, salt with trifluoromethanesulfonic acid (1:1)
(9CI) (CA INDEX NAME)
             CM 1
             CRN 606970-68-7
CMF C17 H25 N2 O2
                          2
                        37181-39-8
C F3 O3 S
```

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
112: 392407
Preparation of monocyclic and bicyclic lactams, in
particular derivatives of pyrrolidines and
pyrroloimidazoles, as factor Xa inhibitors
Han, Wel; Qiao, Jennifer; Hu, Zilun
Bristol-Myers Squibb Company, USA
CODE: PIXXDZ
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:

COPYRIGHT 2006 ACS on STN
2005 ACS on STN
2016 ACS on STN
2017 ACPLIES
PROPAGE ACT NUM. COUNT:
Prepared Information:

COPYRIGHT 2016 ACS on STN
2018 ACS ON STN
2018 ACPLIES
PROPAGE ACT NUM. COUNT:
PIXXD2
Patent INFORMATION:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE PATENT NO. KIND DATE APPLICATION NO. DATE

W0 2005032468 A2 20050414 W0 2004-US31857 20040929

M: AR, AG, ALI, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BR, BC, CA, CRI,
GR, GH, GM, HR, HU, ID, ILI, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
NO, NZ, OM, PG, PH, PH, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW,
AW, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CR, CY, CZ, DE, DK,
EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NLI, PL, PT, RO, SE,
SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GM, GQ, GM, ML, MR, NE,
SN, TD, TG

US 2005107361 A1 20050519 US 2004-952397 20040928

EP 1667647 RI AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PY,
IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, PRIORITY APPLN. INFO.: US 2003-507533P P 20031001 A 20040928 US 2004-952397 WO 2004-US31857 W 20040929

OTHER SOURCE(S): MARPAT 142:392407

```
* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
        Title compds. {I and II; V = (CH2)n; n = 1-3; U = (CH2)m; m = 1-2; one of T1 and T2 = C0, CS, S02, and the other = C0, CS, S02, CH2, CHOH; one of
         and Z2 = N, and the other = C; G = (un)substituted Ph, pyrimidyl,
pyrazinyl, pyridazinyl, etc. optionally fused with a 5-6 membered ring
containing 0-2 heteroatoms; G1 = SO2NH and derivs., NHCO, NHCSNH and
```

containing w-2 necessarians.

(un) substituted alkylene, etc.; A = (un) substituted carbocycle, heterocycle; B = alkylene, SO2H and derivs., (un) substituted carbocyle, heterocycle, etc.; Ria at each occurrence = H, (un) substituted alkylene, alkenylene, alkynylene, etc.; or RiacCRia = (un) substituted 5-7 membered ring; their stereoisomers or pharmaceutically acceptable salts; with

L69 ANSWER 1 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) 503 IT 814254-81-4 RL: CAT (Catalyst use); RCT (Reactant); RACT (Reactant or reagent); USES (Usea) (Uses)

(palladium-carbene catalyzed Sonogashira coupling of unactivated slkyl bromides with alkyl substituted alkynes)
816254-81-4 HCAPLUS
Dispiro[cycloheptane-1,3'(2'H)-imidazo[5,1-b:4,3-b']bisoxazol[4]ium-7'(8'H),1''-cycloheptane), salt with trifluoromethanesulfonic acid (1:1)
(SCI) (CA INDEX NAME) CM 1 CRN 814254-80-3 CMF C19 H29 N2 O2 CM 2 37181-39-8 C F3 O3 S REFERENCE COUNT: THERE ARE 44 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) provisos), were prepd. as inhibitors of trypsin-like serine proteases, specifically Factor Xa. For example, an eleven-step synthesis starting from trans-3-Hydroxy-1-proline is given for lactam III. I displayed Ki \$10 µM for the inhibitors of Factor Xa. I were effective thrombin inhibitors; Ki \$10 µM. I are useful antithrombotics.

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

dimethylaminoethyl]cyclobutyl]phenyl]-1-oxohexahydropyrrolo(1,2-c]imidazol7-yl]amide 85002-54-99, 5-Chlorothiophene-2-carboxylic acid
N-[(7R,7a5)-2-[4-[1-(carbamoylmethyl]cyclobutyl]phenyl]-1oxohexahydropyrrolo(1,2-c]imidazol-7-yl]amide 850002-55-09,
5-Chlorothiophene-2-carboxylic acid
N-[(7R,7a5)-2-[4-(2-dimethylamino-1,1dimethylethyl]phenyl]-1-oxohexahydropyrrolo[1,2-c]imidazol-7-yl]amide
850002-56-19, 5-Chlorothiophene-2-carboxylic acid

N-[(7R,7a5)-1-oxo-2-[4-[1-(pyrrolidin-1-ylmethyl)cyclopropyl)phenyl}hexahy dropyrrolo[1,2-c]imidazol-7-yl]amide 850002-57-2P, 5-Chlorothiophene-2-carboxylic acid N-[(7R,7a5)-2-[4-[1-(morpholin-4-ylmethyl)cyclopropyl]phenyl]-1-oxohexahydropyrrolo[1,2-c]imidazol-7-yl]amide 850002-58-3P, 5-Chlorothiophene-2-carboxylic acid

N-[(7R,7a5)-2-[4-[1-[(5-methylthiazol-2-ylamino)methyl]cyclopropyl]phenyl]-l-oxohexahydropyrcolo[1,2-c]imidazol-7-yl]amide
RI: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (III.e.)

(drug candidate; prepn. of monocyclic and bicyclic lactams as Factor Χa

inhibitors)
850001-02-4 HCAPLUS
2-Thiophenecarboxamide, 5-chloro-N-[(7R,7aS)-hexahydro-1-oxo-2-[4-{2-oxo-1(2H)-pyridinyl)phenyl]-1H-pyrrolo[1,2-c]imidazol-7-yl]- (9CI) (CA INDEX MANEY)

Absolute stereochemistry.

RN 850001-03-5 HCAPLUS
CN -2-Thiophenecarboxamide,
5-chloro-N-(7R,7a5)-hexahydro-1-oxo-2-[4-(2-oxo-1-piperidiny1)pheny1]-1H-pyrrolo[1,2-c]imidazo1-7-y1]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

N 850002-06-1 HCAPLUS N 2-Thiophenecarboxamide, -chloro-N-(17R,7aS)-hexahydro-1-oxo-2-{4-{1-{2-(1-pyrrolidiny)}ethy]eyclobutyl]phenyl]-1H-pyrrolo[1,2-c]imidazol-7-yl]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

850002-07-2 HCAPLUS
2-Thiophenecarboxamide, 5-chloro-N-[(7R,7aS)-hexahydro-2-[4-[1-(4-

morpholinylmethyl)cyclobutyl]phenyl]-1-oxo-1H-pyrrolo[1,2-c]imidazol-7-yl](9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 850001-06-8 HCAPLUS
CN 1H-Indole-6-carboxamide,
3-chloro-N-[(7R,7a8]-hexahydro-1-oxo-2-[4-(2-oxo1(2K)-pyridinyl)phenyl]-1H-pyrrolo[1,2-c]imidazol-7-yl]- (9CI) (CA INDEX

Absolute stereochemistry.

RN 850001-07-9 HCAPLUS
CN 1H-Indole-6-carboxamide,
3-chloro-M-([R, 7a5]-hexahydro-1-oxo-2-[4-(3-oxo-4-morpholinyl)phenyl]-1H-pyrrolo[1,2-c]imidazo1-7-yl]- (9CI) (CA INDEX

Absolute stereochemistry.

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
RN 850002-08-3 HCAPLUS
CN 2-Thiophenecarboxamide, 5-chloro-N-[(7R,7aS)-hexahydro-2-[4-[1-[2-(4-

Absolute stereochemistry.

850002-09-4 HCAPLUS
2-Thiophenecarboxamide, 5-chloro-N-{(7R,7aS)-2-{4-{1-{2-}}}
(dimethylamino)ethyl]cyclopentyl]phenyl]hexahydro-1-oxo-1H-pyrrolo[1,2-c]imidarol-7-yl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 850002-10-7 HCAPLUS
CN 2-Thiophenecarboxamide,
5-chloro-M-(1R,7a8)-hexahydro-1-oxo-2-[4-{1-[2-(1pyrrolidinyl)ethyl]cyclopropyl]phenyl]-1H-pyrrolo[1,2-c]imidazo1-7-yl](SCI) (CA INDEX NAME)

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

850002-11-8 HCAPLUS
2-Thiophenecarboxamide, 5-chloro-N-[(7R,7aS)-hexahydro-2-[4-[1-[2-(4-morpholiny]) ethyl]-cyclopropyl]phenyl]-1-oxo-1H-pyrrolo[1,2-c]imidazol-7-yl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

850002-12-9 HCAPLUS 2-Thiophenecarboxamide, 5-chloro-N-[(7R,7aS)-2-[4-[1,1-dimethyl-3-(1-

Absolute stereochemistry.

ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) 850002-15-2 HCAPLUS 2-Thiophenecarboxamide, 5-chloro-N-[{7R,7aS}-hexahydro-2-[4-[1-(methoxymethyl)cyclobutyl]phenyl]-1-oxo-lH-pyrrolo[1,2-c]imidazol-7-yl]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

850002-47-0 HCAPLUS
2-Thiophenecarboxamide, 5-chloro-N-[(7R,7aS)-2-[4-[1-(dimethylamino)cyclopropyl]phenyl]hexahydro-1-oxo-lH-pyrrolo[1,2-c]imidazol-7-yl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

850002-48-1 HCAPLUS
2-Thiophenecarboxamide, 5-chloro-N-[(7R,7as)-2-[4-[1-[(dimethyl]amino]methyl]cyclopropyl]phenyl]hexahydro-1-oxo-1H-pyrrolo[1,2-c]imidazol-7-yl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

850002-13-0 HCAPLUS
2-Thiophenecarboxamide, 5-chloro-N-{(7R,7aS}-2-{4-{1,1-dimethyl-3-(4-

morpholinyl)propyl]phenyl]hexahydro-1-oxo-1H-pyrrolo[1,2-c]imidazol-7-yl][9CI] (CA INDEX NAME)

Absolute stereochemistry.

850002-14-1 HCAPLUS
2-Thiophenecarboxamide, 5-chloro-N-{{7R,7aS}-hexahydro-2-[4-[1-(methoxymethyl]cyclopropyl]phenyl}-1-oxo-1H-pyrrolo[1,2-c]imidazol-7-yl]-(SCI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

850002-49-2 HCAPLUS
2-Thiophenecarboxamide, 5-chloro-N-[(7R,7as)-2-[4-[1-[2-(dimeth)lamino)ethyl]cyclopropyl]phenyl]hexahydro-1-oxo-1H-pyrrolo[1,2-c]imidazol-7-yl]- (9CI) (CA INDEX NAME)

850002-50-5 HCAPLUS
2-Thiophenecarboxamide, N-{{7R,7as}-2-{4-{1-(2-amino-2-oxoethyl-leyclopropyl]phenyl]hexahydro-1-oxo-1H-pyrrolo{1,2-c}imidazol-7-yl}-5-chloro-{9C1} (CA INDEX NAME)

Absolute stereochemistry.

850002-51-6 HCAPLUS 2-Thiophenecarboxamide, 5-chloro-N-[(7R,7as)-2-[4-[1-

(dimethylamino)cyclobutyl]phenyl]hexahydro-1-oxo-1H-pyrrolo{1,2-c}imidazol-7-yl]- (9CI) (CA INDEX NAME)

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 850002-52-7 HCAPLUS
CN 2-Thiophenecarboxamide, 5-chloro-N-[(7R,7as)-2-[4-[1[(dimethylamino)methyl]-cyclobutyl|phenyl]hexahydro-1-oxo-1H-pyrrolo[1,2c]imidazol-7-yl}- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 850002-53-8 HCAPLUS
CN 2-Thiophenecarboxamide, 5-chloro-N-[(7R,7aS)-2-{4-[1-[2-(dimethylamino)ethyl]cyclobutyl]phenyl]hexahydro-1-oxo-1H-pyrrolo[1,2-c]imidazol-7-yl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 850002-54-9 HCAPLUS

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 850002-57-2 HCAPLUS
CN 2-Thiophenecarboxamide, 5-chloro-N-[(7R,7aS)-hexahydro-2-[4-[1-(4-morpholinylmethyl)cyclopropyl]phenyl]-1-oxo-1H-pyrrolo[1,2-c]imidazol-7-yl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 850002-58-3 HCAPLUS
CN 2-Thiophenecarboxamide,
5-chloro-N-[(7R,7as)-hexahydro-2-[4-[1-[(5-methyl2-thiazolyl)lamino]methyl]cyclopropyl]phenyl]-1-oxo-1H-pyrrolo[1,2c)imidazol-7-yl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) CN 2-Thiophenecarboxamide, N-[(7R,7as]-2-[4-[1-(2-amino-2-

oxoethyl)cyclobutyl]phenyl]hexahydro-1-oxo-1H-pyrrolo[1,2-c]imidazol-7-yl]-5-chloro- [9CI] (CA INDEX NAME)

Absolute stereochemistry.

RN 850002-55-0 HCAPLUS
CN 2-Thiophenecarboxamide, 5-chloro-N-[(7R,7as)-2-(4-[2-[dimethylamino)-1,1-dimethylethyl] phenyl | hexahydro-1-oxo-1H-pyrrolo[1,2-c]imidazo1-7-yl]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 850002-56-1 HCAPLUS
CN 2-Thiophenecarboxamide, 5-chloro-N-{(7R,7as}-hexahydro-1-oxo-2-[4-[1-{1-pyrcolidinylmethyl]cyclopropyl]phenyl]-1H-pyrrolo[1,2-c]imidazo1-7-yl](9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 2 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L69 ANSWER 3 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1171LE:
12:176629
Organocatalyzed conjugate umpolung of o, β-unsaturated aldehydes for the synthesis of γ-butyrolactones
AUTHOR(S):
Burstein, Christian Glorius, Frank
MAX-Planck-Institut Lucr Kohlenforshung, Muelheim an der Ruhr, 45470, Germany
Angewandte Chemie, International Edition (2004), 43(45), 6205-6208
CODEN: ACIEFS: ISSN: 1433-7851
Wiley-VCH Verlag GmbH & Co. KGaA
JOURNAL
LANGUAGE:
OTHER SOURCE(S):
CASREACT 142:176629

OTHER SOURCE(S):

N-heterocyclic carbenes can generate homoenolate equivalent under mild conditions by conjugate umpolung of  $\alpha,\beta$ -unsatd. aldehydes. This organocatalytic reaction allows an efficient one-step synthesis of substituted  $\gamma$ -butyrolactones. E.g., the N-heterocyclic carbene generated from imidazolium I was used to catalyze the reaction of (E)-PhCH:CHCHO with 4-ClC6H4CHO to give 53%  $\gamma$ -butyrolactone II (80:20 cia/trans). 832098-68-7

CM 1

L69 ANSWER 4 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:964820 HCAPLUS
DOCUMENT NUMBER: 141:395584
Preparation of novel triazine compounds for inhibiting

INVENTOR (S):

rreparation of novel triazine compounds for smooth muscle cell proliferation Timmer, Richard T.; Alexander, Christopher W.; Pillarisetti, Sivaram: Saxena, Uday; Yeleawarapu, Koteswar Rao: Pal, Manojlt: Reddy, Jangalgar Tirupathy: Reddy, Velagala Venkira Rama Murali Krishna: Sridevi, Bhatlapenumarphy Shesha; Kumar, Potlapally Rajender; Reddy, Gaddam Om USA USA: U.S. Pat. Appl. Publ., 433 pp., Cont.-in-part of U.S. Ser. No. 390,485.
CODEN: USXXCO Patent English 6

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. PATENT NO. DATE KIND US 2004224950 US 2004077648 PRIORITY APPLN. INFO.: 20041111 US 2003-400140 US 2003-390485 20030326 A1 A1 US 2001-324147P US 2002-253388 B1 20020923 US 2003-390485 A2 20030317

OTHER SOURCE(S): MARPAT 141:395584

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

The present invention relates to methods and compns. comprising compds. I or II [Rl = H, alkyl, cycloalkyl, etc.; G = NRl, O; J = CH, N; n = 0-3;

= o-Rl, m-Rl, m-ORl, m-OCF3, etc.; X2 = o-Rl, p-Rl, p-ORl, p-OCF3, etc.; X3 = o-Rl, m-Rl, p-Rl, o-ORl, p-ORl; or X2 and X3 together is a fused benzene, pyridine, dioxane, tetrahydropyran ring; AY, DY = ORl, F, Cl,

I, tetrahydroquinolin-1-yl, etc.; or A, B = 0, NR1; and Y = R1,

[CHR1]qCI, (CHR1)qCI3, etc.; q = 0-3] that treat pathophysiol. conditions arising from inflammatory responses. Over 100 synthetic examples described synthesis of compds. I and II and their intermediates. E.g., a

synthesis of compus. a since it is not constructed by the synthesis of the triazine III, starting from cyanuric chloride, is given. In particular, the present invention is directed to compus. that inhibit or block glycated protein produced induction of the signaling-associated inflammatory response in endothelial cells. The present invention

relates
to compds. that inhibit smooth muscle cell (SMC) proliferation. Many of
the compds. I and II inhibited SMC proliferation by greater than 70%.
Also, the most effective compds. I and II showed an 80% decrease in II-6

L69 ANSWER 3 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN CRN 832098-67-6 CMF C12 H19 N2 O2 (Continued)

Absolute stereochemistry.

2 CM

CRN 37181-39-8 CMF C F3 03 S

REFERENCE COUNT:

THERE ARE 56 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L69 ANSWER 4 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) secretion in test for AGE-induced inflammatory response detn. In particular, the present invention is directed to compds. that inhibit smooth muscle cell proliferation by modulating HSPGs such as Perlecan. The present invention further relates to the use of compds. to treat vascular occlusive conditions characterized by smooth muscle proliferation such as restenosis and atherosclerosis.

IT 676358-28-49 676358-97-7P
R1: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study): PREP (Preparation); USES (Uses)

(preparation of novel triazine compds. for inhibiting smooth muscle

(preparation of novel triazine compds. for inhibiting smooth muscle

proliferation)
676358-28-4 HCAPLUS
1,3,5-Triazine-2,4-diamine, N-(cyclohexylmethyl)-N'-(3-fluoro-4-methoxyphenyl)-6-(tetrahydro-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)- (9CI)
(CA INDEX NAME)

676358-97-7 HCAPLUS
1H-Pyrrolo(1,2-c)imidazol-1-one, 2-[4-[(cyclohexylmethyl)amino]-6-[(3-fluoro-4-methoxyphenyl)amino]-1,3,5-triazin-2-yl]hexahydro- (9CI) (CA INDEX NAME)

L69 ANSWER 4 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) L69 ANSWER 5 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:902612 HCAPLUS DOCUMENT NUMBER: 142:93738 2004:902612 HCAPINS
Sterically demanding, bioxabiline-derived
N-heterocyclic carbene ligands with restricted
flexibility for catalysis
Altenhoff, Geroon; Goddard, Richard; Lehmann,
Christian W.; Glorius, Frank
Max-Planck-Institut fuer Kohlerforschung, Muelheim an
der Ruhr, 45470, Germany
Journal of the American Chemical Society (2004)
126(46, 15195-15201
CODEN: JASSAT; ISSNT 0002-7863
American Chemical Society
Journal
Journal DOCUMENT NUMBER: TITLE: AUTHOR (S): CORPORATE SOURCE: SOURCE: PUBLISHER: DOCUMENT TYPE: LANGUAGE: Journal English OTHER SOURCE (S): CASREACT 142:93738

The triflate salts of imidazobioxazolium ions I [R = Rl = Me; RRl = (CH2)n; n = 5, 6, 7, 8, 12] are prepared as precursors for sterically demanding and conformationally constrained N-heterocyclic carbene (NNC) ligands; palladium complexes derived from I [RRl = (CH2)n; n = 7, 12] act as effective catalysts for the Suzuki-Miyaura coupling reactions of ortho-substituted aryl chlorides with ortho-substituted arylboronic acids to provide triortho- and tetraortho-substituted biaryls such as II in 47-96% yields. I=CF3SO3- are prepared in five steps from on,-disubstituted amino acids and di-Et oxalate; reduction of amino acids to the amino acids to the amino alcs., condensation of the amino alcs. with

di-Et coxalate to give the hydroxymethyl-substituted oxamides, chlorination of the primary alc. moleties, cyclization of the oxamide with the chloromethyl groups to give the bioxazolines, and reaction of the bioxazolines with chloromethyl pivalate and silver triflate. IscT3803- are soluble in methylene chloride and THF and are chromatographable. Iridium cyclooctadienyl and iridium dicarbonyl chloride complexes derived from IsCT3803- [R = Rl = Me; RRl = (CH2)n; n = 6, 8 , 12) are prepared; IR frequencies of the carbonyl ligands indicate

That carbene ligands derived from I=CF3803- are less electron-donating than previous NHC ligands but are comparable to electron-rich phosphines. Selected iridium cyclooctadienyl and iridium dicarbonyl chloride

of imidazobioxazolium ligands are characterized by X-ray crystallog.
Dimeric palladium chloride complexes derived from I=CF3SO3- (RR1 =

ANSWER 5 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) (CH2)n; n = 7, 12] are prepd. and characterized by X-ray crystallog. Generation of the carbene ligand from I=CF3SO3- [RR1 = (CH2)12] by treatment with potassium hydride and potassium tert-butoxide followed by addn. of palladium acetate yields a palladium catalyst which is effective for the Suzuki-Miyaura coupling of highly hindered aryl chlorides and arylboronic acids. Potassium phosphate is the most effective base and toluene is the most effective solvent for Suzuki-Miyaura coupling of highly hindered aryl chlorides and arylboronic acids using imidazobioxazolium-derived carbene ligands, although cesium carbonate can also be used as the base and 1,4-dioxane as the solvent; the isolated dimeric paladium chloride complexes derived from I=CF3SO3- [RR1 = (CH2)12] can also be used as catalysts. Anhyd. conditions are important to minimize hydrodeborylation byproducts of the coupling reaction. E.g., in the presence of the palladium catalyst generated from I=CF3SO3- [RR1 = (CH2)12] and palladium acetate and potassium phosphate, 2-chloro-1, 3-dimethylbenzene and 2,4,6-trimethylphenylboronic acid undergo coupling in toluene at 100° for 16 h to provide biphenyl I in 96% yield.
814254-81-49 814254-83-69
RR: CAT (Catalyst use): PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (preparation and crystal structure of an imidazobioxazolium triflate its IT

use as a precursor for a sterically hindered and electron-donating N-heterocyclic carbene ligand in Suzuki-Miyaura coupling reactions of hindered aryl chlorides)
814254-81-4 HCAPLUS
Dispiro[cycloheptane-1,3'(2'H)-imidazo[5,1-b:4,3-b']bisoxazol[4]ium-7'(8'H),1''-cycloheptane], salt with trifluoromethanesulfonic acid (1:1)
(9CI) (CA INDEX NAME)

CM 1 CRN 814254-80-3 CMF C19 H29 N2 O2

CM 2 CRN 37181-39-8 CMF C F3 03 S

L69 ANSWER 5 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN 814254-83-6 HCAPLUS
Dispiro[cyclooctane-1,3'(2'H)-imidazo[5,1-b:4,3-b']bisoxazol[4]ium-7'(8'H),1''-cyclooctane], salt with trifluoromethanesulfonic acid (1:1)
(9CI) (CA INDEX NAME) CM 1 CRN 814254-82-5 CMF C21 H33 N2 O2

CRN 37181-39-8 CMF C F3 O3 S

CM 2

F-C-503

IT 814254-79-0P RE: CAT (Catalyst use); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); USES (Uses) (preparation and crystal structure of an imidazobioxazolium triflate

and its use as a precursor for a sterically hindered and electron-donating N-heterocyclic carbene ligand in Suzuki-Miyaura coupling reactions of hindered aryl chlorides) 814254-79-0 MCAPUS

Dispiro[cyclopentane-1,3'(2'H)-imidazo[5,1-b:4,3-b']bisoxazol[4]lum-7'(0'H),1''-cyclopentane), salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1 CRN 814254-78-9 CMF C15 H21 N2 O2 L69 ANSWER 5 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

606970-69-89
RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and crystal structure of an imidazobioxazolium triflate

use as a precursor for a sterically hindered and electron-donating N-heterocyclic carbene ligand in Suzuki-Miyaura coupling reactions of hindered aryl chlorides) (606970-69-8 HCAPLUS Dispiro[cyclohexane-1,3'(2'H)-imidazo[5,1-b:4,3-b']bisoxazol[4]ium-7'(8'H],1''-cyclohexane], salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 606970-68-7 CMF C17 H25 N2 O2

CM 2

CRN 37181-39-8

L69 ANSWER 5 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

CRN 37181-39-8 CMF C F3 O3 S

814254-85-8P

814236-85-89
RL: CAT (Catalyst use); RCT (Reactant); SPN (Synthetic preparation); PREP
(Preparation); RACT (Reactant or reagent); USES (Uses)
 (preparation of an imidazobioxazolium triflate and its use as a

or for a sterically hindered and electron-donating N-heterocyclic carbene ligand in Suzuki-Miyaura coupling reactions of hindered aryl

chlorides) RN 814254-85-8 HCAPLUS

Dispiro(cycloddecane-1,3'(2'H)-imidszo[5,1-b:4,3-b']bisoxazol[4]ium-7'(8'H),1''-cycloddecane], salt with trifluoromethanesulfonic acid (1:1)(9CI) (CA INDEX NAME)

CM 1

CRN 814254-84-7 CMF C29 H49 N2 O2

CM

RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP

L69 ANSWER 5 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN CMF C F3 O3 S (Continued)

CM 1

814254-86-9
RL: PRP (Properties)
(preparation and crystal structure of an imidazobioxazolium triflate precursor for a sterically hindered and electron-donating N-heterocyclic carbene ligand)
814254-86-9 HCAPLUS
Dispiro[cyclododecane-1, 3' (2'H)-imidazo[5, 1-b:4, 3-b']bisoxazol[4]ium-7'(8'H), 1''-cyclododecane], salt with trifluoromethanesuifonic acid, compd. with dichloromethane (1:1:1) (9CI) (CA INDEX NAME)

CRN 75-09-2 CMF C H2 C12

C1-CH2-C1

CM 2

CRN 814254-85-8 CMF C29 H49 N2 O2 . C F3 O3 S

CM 3

CRN 814254-84-7 CMF C29 H49 N2 O2

L69 ANSWER 5 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
(Preparation): RACT (Reactant or reagent)
(prepn. of imidazobioxazolium triflates as precursors for sterically
hindered and electron-donating N-heterocyclic carbene ligands)
RN 814254-77-8 HCAPLUS
CN Imidazo(5,1-b:4,3-b')bisoxazol-4-ium, 2,3,7,8-tetrahydro-3,3,7,7tetramethyl-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA
INDEX NAME)

CM 1

2 CM

REFERENCE COUNT: THIS

THERE ARE 67 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L69 ANSWER 6 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 2004:878154 HCAPLUS COPYRIGHT 2016 ACS ON STN 2004:878154 HCAPLUS 11TILE: 11TIL

Preparation of novel triazine compounds for

INVENTOR (S)

smooth muscle cell proliferation
Timmer, Richard T.; Alexander, Christopher W.;
Pillarisetti, Sivaram: Saxena, Uday; Yeleswarapu,
Koteswar Rao: Pal, Manojit; Reddy, Jangalgar
Tirupathy; Krishma, Reddy Velagala Venkata Rama
Murali; Sesila, Sridevi Bhatlapenumarthy; Kumar,
Potlapally Rajender; Reddy, Gaddam Om
USA
U.S. Pat. Appl. Publ., 422 pp.
CODEN: USXXCO
Patent
English
6

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE APPLICATION NO. KIND DATE US 2004209882 US 2005124619 PRIORITY APPLN. INFO.: 20041021 US 2003-400169 US 2004-951120 US 2001-324147P 20030326 20040927 20010921 A2 20030317 US 2003-390485 A3 20030326 US 2003-400169

OTHER SOURCE(S): MARPAT 141:366254

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

The present invention relates to methods and compns. comprising compds. I or II  $\{Rl=H,\ alkyl,\ cycloalkyl,\ etc.;\ G\approx NRl,\ O;\ J=CH,\ N;\ n=0-3;$ 

= o-R1, m-R1, m-OR1, m-OCF3, etc.; X2 = o-R1, p-R1, p-OR1, p-OCF3, etc.; X3 = o-R1, m-R1, p-R1, o-OR1, p-OR1; or X2 and X3 together is a fused benzene, pyridine, dioxane, tetrahydropyran ring; AY, DY = OR1, F, C1,

I, tetrahydroquinolin-1-yl, etc.; or A, B = O, NR1; and Y = R1,

I, tetranygroquinozin-s-ys, conditions arising (CHR1)qR1, (CHR1)qCF3, etc.; q = 0-3) that treat pathophysiol. conditions arising from inflammatory responses. Over 100 synthetic examples described synthesis of compds. I and II and their intermediates. E.g., a multi-sten

I-step synthesis of the triazine III, starting from cyanuric chloride, is given In particular, the present invention is directed to compds. that inhibit or block glycated protein produced induction of the signaling-associated

L69 ANSWER 6 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L69 ANSWER 6 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) inflammatory response in endothelial cells. The present invention

inflammatory response in endothelial cells. The present invention relates to compds. that inhibit smooth muscle cell (SMC) proliferation. Many of the compds. I and II inhibited SMC proliferation by greater than 70%. Also, the most effective compds. I and II showed an 80% decrease in IL-6 secretion in test for AGE-induced inflammatory response detn. In particular, the present invention is directed to compds. that inhibit smooth muscle cell proliferation by modulating MSPGs such as Perlecan. The present invention further relates to the use of compds. to treat vascular occlusive conditions characterized by smooth muscle proliferation such as restenosis and atherosclerosis.

If \$76358-28-4P 676358-97-7P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of novel triazine compds. for inhibiting smooth muscle

proliferation)
1,3,5-Triazine-2,4-diamine, N-(cyclohexylmethyl)-N'-(3-fluoro-4-methoxyphenyl)-6-(tetrahydro-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)- (9CI)
(CA INDEX NAME)

676358-97-7 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-l-one, 2-[4-[(cyclohexylmethyl)amino]-6-[(3-fluoro-4-methoxyphenyl)amino]-1,3,5-triazin-2-yl]hexahydro- (9CI) (CA INDEX NAME)

L69 ANSWER 7 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:878153 HCAPLUS DOCUMENT NUMBER: 141:366253 TITLE: preparation of novel triazine cinhibiting Preparation of novel triazine compounds for

smooth muscle cell proliferation
Timmer, Richard T.; Alexander, Christopher W.;
Pillarisetti, Sivaram; Saxena, Uday; Yeleswarapu,
Koteswar Rao; Pal, Manojit; Reddy, Jangalgar
Tirupathy; Krishna, Reddy Velagala Venkata Rama
Murali; Sridevi, Bhatlapenumarthy Sesha; Kumar,
Potlapally Rajender; Reddy, Gaddam Om
USA
U.S. Pat. Appl. Publ., 254 pp.
CODEN: USXXCO
Patent
English
6 INVENTOR (S):

PATENT ASSIGNEE (S):

SOURCE:

DOCUMENT TYPE: LANGUAGE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE US 2003-400134 US 2004-951305 US 2001-324147P 20030326 20040927 P 20010921 US 2004209881 US 2005113341 PRIORITY APPLN. INFO.: 20041021 B1 20020923 A2 20030317

US 2003-400134

OTHER SOURCE(S): MARPAT 141:366253

The present invention relates to methods and compns. comprising compds. I  $\{Rlb = substituted\ Ph;\ R2b = 1-indolyl,\ substituted\ NH2,\ substituted$ 

substituted OH, etc.; R6b = O, NH, NMe, NEt, N(CN); R7b = cycloheptanyloxy, cyclopropanyloxy, cyclopentanyloxy, cyclohexanyloxy, substituted NH2] that treat pathophysiol. conditions arising from inflammatory responses. Over 100 synthetic examples described synthesis of compds. I and their intermediates. E.g., a multi-step synthesis of

A3 20030326

L69 ANSWER 7 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) triazine II, starting from cyanuric chloride, is given. In particular, the present invention is directed to compds. that inhibit or block glycated protein produced induction of the signaling-associ inflammatory response in endothelial cells. The present invention relates to compds. that inhibit smooth muscle cell [SMC] proliferation. Many of the compds. I inhibited SMC proliferation by greater than 70s. In particular, the present invention is directed to compds. that inhibit smooth muscle cell proliferation by modulating HSPGs such as Perlecan. The present invention further relates to the use of compds. to treat vascular occlusive conditions characterized by smooth muscle proliferation such as restenosis and atherosclerosis.

676357-62-3P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation) of novel triazine compds. for inhibiting smooth muscle cell

cell
proliferation)
RN 676357-62-3 HCAPLUS
CN 1,3,5-Triazine-2,4-diamine,
N-cycloheptyl-N'-(3-fluoro-4-methoxyphenyl)-6(tetrahydro-iH-pyrrolo[1,2-c]imidarol-2(3H)-yl)- (9CI) (CA INDEX NAME)

L69 ANSWER 8 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:267312 HCAPLUS DOCUMENT NUMBER: 140:303704 TITLE: Preparation of aminotriazines

140:303704
Preparation of aminotriazines for treatment of unvanted cell proliferation, inflammation, hyperproliferation, and as glycosidase modulators. Timmer, Richard T., Alexander, Christopher W.; Pillarisetti. Sivaram: Saxena, Uday; Yeleswarapu, Koteswar Rao: Pal, Manopit: Reddy, Jangalgar Tirupathy: Reddy, Velagala Venkata Rama Murali Krishna; Sridevi, Bhatlapenumarthy Sesha; Kumar, Potlapally Rajender: Reddy, Gaddam Om Reddy US Therapeutics, Inc., USA PCT Int. Appl., 840 pp.
CODEN: PIXXD2
Patent

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: English 6

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

INVENTOR(S):

	PA1	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D.	ATE	
		2004									WO 2	003-	US93	56		2	0030	326
	WO	2004																
		W:										BG,						
												EΕ,						
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KĢ,	KP,	KR,	ΚZ,	LC,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	ΜK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,	OM,
			PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	sĸ,	SL,	ΤJ,	TM,	TN,	TR,	TT,
												ZM,						
		RW:	GH,	GM,	ΚĒ,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
			KG,	KZ,	MD,	RU,	TJ,	TM.	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
			FI,	FR,	GB,	GR,	ΗU,	IE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,
												G₩,						
	US	2004	0776	48		A1		2004	0422		US 2	003-	3904	85		2	0030	317
	CA	2499	964			AA		2004	0401		CA 2	003-	2499	964		2	0030	326
	ΑŲ	2003	2319	75		A1		2004	0408		AU 2	003-	2319	75		2	0030	326
	₿R	2003	0146	70		A		2005	0809		BR 2	003-	1467	0		2	0030	326
	EP	1560	817			A1		2005	0810		EP 2	003-	7977	88		2	0030	326
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	ΗU,	SK	
	JP	2006	5114	76		T2		2006	0406		JP 2	004-	5381	53		2	0030	326
PRIOF	IT	APP	LN.	INFO	.:						US 2	002-	2533	88		A 2	0020	923
											US 2	003-	3904	85		A 2	0030	317
											US 2	001-	3241	47P		₽ 2	0010	921
												003-						

OTHER SOURCE(S): MARPAT 140:303704

L69 ANSWER 8 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

Title compds. e.g. [I: Rl = substituted Ph. PhCH2, PhCH2CH2, pyridyl; R2

(substituted) amino, piperazinyl, piperidinyl, thiomorpholinyl, piperidinylamino, hydroxymethylpyrrolidinyl; R6 = H, Me; R7 = hexamethyleneimino, cycloheptylimino, bicyclo[2.2.1]heptyloxy,

substituted amino], were prepared Thus,

amino], were prepared Thus,

N2-(3-chloro-4-methoxyphenyl)-N4-cycloheptyl-N6methyl-N6-piperidin-4-yl-1, 3,5-triazine-2,4,6-triamine in an
antiproliferation assay (perlican) showed ICSO = 2.2 µM.

IT 676357-62-3P, N-Cycloheptyl-N'-(3-fluoro-4-methoxyphenyl)-6(tetrahydropyrrolo[1,2-c]imidazol-2-yl)-[1,3,5]triazine-2,4-diamine
676358-28-4P, N-Cyclohexylmethyl-N'-(3-fluoro-4-methoxyphenyl)-6(tetrahydropyrrolo[1,2-c]imidazol-2-yl)-[1,3,5]triazine-2,4-diamine
676358-97-7P, 2-[4-(Cyclohexylmethylamino)-6-(3-fluoro-4-

methoxyphenylamino) - [1,3,5] triazin-2-yl] hexahydropyrrolo[1,2-c]imidazol-1-

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uses)

(Uses)

(preparation of aminotriazines for treatment of unwanted cell proliferation,

inflammation, hyperproliferation, and as glycosidase modulators)

RN 67637-62-3 HCAPLUS

CN 1,3,5-Triazine-2,4-diamine,

N-cycloheptyl-N'-(3-fluoro-4-methoxyphenyl)-6
(tetrahydro-1H-pyrrolo[1,2-c]imidarol-2(3H)-yl)- (9CI) (CA INDEX NAME)

676358-28-4 HCAPLUS
1,3,5-Triazine-2,4-diamine, N-{cyclohexylmethyl}-N'-(3-fluoro-4-methoxyphenyl)-6-{tetrahydro-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)- (9CI) (CA INDEX NAME)

L69 ANSWER 8 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

676358-97-7 HCAPLUS
1H-Pyrrolo[1,2-c|imidazol-1-one, 2-[4-[(cyclohexylmethyl)amino]-6-[(3-fluoro-4-methoxyphenyl)amino]-1,3,5-triazin-2-yl]hexahydro- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

(Continued)

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

CRN 606970-68-7 CMF C17 H25 N2 O2

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L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER: 2004:60481 HCAPLUS
TITLE: 140:128420
preparation of imidazolium salts from bisimines and alkylating agents in the presence of metal salts as promoters.
Glorius, Frank
Studiengesellschaft Kohle mbH, Germany
PCT Int. Appl.. 35 pp.
CODEN: PIXXD2
PATENT TYPE: PETAL PROMOTE PIXXD2
PATENT INFORMATION: 1
     DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                 PATENT NO.
                                                                                                                                                   KIND
                                                                                                                                                                                      DATE
                                                                                                                                                                                                                                                             APPLICATION NO.
PATENT NO. KIND DATE

2004017465

W: RE, AL, AM, AT, AU, AZ, BA,
CR, CU, C2, DK, DM, D2, EC,
HU, ID, II, IN, IS, JP, KE,
LU, LV, MA, MD, MG, KK, MN,
RO, RU, SC, SD, SE, SG, SK,
US, UZ, VC, VN, YU, 2A,
RW: GH, GM, KE, LS, MW, MZ, SD,
KG, KZ, MD, RU, TJ, TM, AT,
FI, FR, GB, GR, HU, IE, IT,
BF, BJ, CF, CG, CI, CM, GA,
DE 10231368

AU 2003247251

AU 2003247251

R: AT, BE, CH, DE, DK, ES, FR,
IE, SI, LT, LV, FI, RO, MK,
JP 2005538071

US 2005240025

PRIORITY APPLN. INFO:
                                                                                                                                                                                                                                       APPLICATION NO.

***O 2003-DE2285
BB, BG, BR, BY, BZ, EE, ES, FI, GB, GD, KG, KP, KR, KZ, LC, MM, MK, MZ, NO, MZ, SL, TJ, TM, TM, TR, ZW
SL, SZ, TZ, UG, ZM, BE, BG, CH, CY, CZ, LU, MC, NL, PT, RO, GN, GO, GM, ML, RR, DE 2002-10231368
AU 2003-247251
EP_2003-105451
GB, GR, TT, LI, LY, AL, TR, BG, CZ, JP 2004-520311
US 2005-520800
DE 2002-10231368
                                                                                                                                                                                                                                                                                                                                                                                 20030708
CH, CN, CO,
GH, GM, HR,
LR, LS, LT,
PH, PL, PT,
T2, UA, UG,
                                                                                                                                                                                                                                                                                                                                                                               AM, AZ, BY,
DK, EE, ES,
SI, SK, TR,
SN, TD, TG
20020711
20030708
SE, MC, PT,
HU, SK
20030708
20050110
A 20020711
                                                                                                                                                                                                                                                             WO 2003-DE2285
                                                                                                                                                                                                                                                                                                                                                                               w 20030708
    OTHER SOURCE(S):
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MARPAT 140:128420

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AB Title compds. [I, II, III; R1-R14 = (unsatd.) (substituted) (cyclic) alkyl, alkenyl, alkynyl, aralkyl, aryl; R1-R8, R11, R13 may addnl. = H; R11, R13 may addnl. = OR16, SR17, NR18R19; R16-R19 = R1; R1, R2, R7, R8, R12, R14, R16-R19 can = linker to another imidazolium residue; X, Y = O, S, (aubstituted) imino; A = (in)organic (polyvalent) anion] were prepared by reaction of the corresponding bisimines with ZCH2OCO2R15, ZCH2O2CR15, or ZCH2OR15 (Z = leaving group; R15 = R3) in the presence of MR (M = (polyvalent) metal cation, tetraorganoammonium, triorganosilyi; A as above). Thus, AgOTf and ClCH2O2CCMe3 were stirred 45 min. in CH2C12; the resulting solution was added to bisoxazoline (IV) followed by stirring for 20 h at 40° to give 85% title commound (V). V was used as a cocatalvate.
for 20

h at 40° to give 85% title compound (V). V was used as a cocatalyst in Suzuki coupling reactions using sterically hindered aryl chlorides.

IT 606970-69-8P

RL: CAT (Catalyst use); IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)
(preparation of imidazolium salts from bisimines and alkylating agents in
the presence of metal salts as promoters)

RN 606970-69-8 HCAPLUS
CN Dispiro(cyclohexane-1,3'(2'H)-imidazo(5,1-b:4,3-b')bisoxazol(4)ium-7'(8'H),1''-cyclohexane], salt with trifluoromethanesulfonic acid (1:1)
(9CI) (CA INDEX NAME)
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L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) CM 2 CRN 37181-39-8 CMF C F3 O3 S - 503 512193-98-5P 512194-01-3P 512194-04-6P
648929-69-1P 648929-51-5P 648929-53-TP
648929-57-1P 648929-55-3P 648929-61-TP
648929-63-PP 648929-55-1P 648929-67-3P
648929-67-5P 648929-71-5P 648929-73-1P
648929-15-3P 648929-71-5P 648929-73-1P
648929-91-1P 648929-83-3P 648929-91-3P
648929-91-7P 648929-89-3P 648929-91-3P
648929-91-7P 648929-89-7P 648929-97-9P
648929-99-1P 648930-01-2P 648930-01-3P
648930-12-5P 648930-74-7P 648930-10-3P
648930-12-5P 648930-14-7P 648930-10-3P
648930-14-1P
648930-18-1P
64893 IT ts in the presence of metal salts as promoters)
512193-98-5 HCAPLUS
Imidazo[5,1-b:4,3-b']bisoxszol-4-ium, 2,3,7,8-tetrahydro-3,7-bis(1-methylethyl)-, (38,75)-, salt with trifluoromethanesulfonic acid (1:1)
(9CI) (GA INDEX NAME) CM 1 CRN 512193-97-4 CMF C13 H21 N2 O2 Absolute stereochemistry. Rotation (+).

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) CM 2 503 512194-01-3 HCAPLUS Imidazo[5,1-b:4,3-b']bisoxazol-4-ium, 3,7-bis(1,1-dimethylethyl)-2,3,7,8-tetrahydro-, (3S,7S)-, salt with trifluoromethanesulfonic acid (1:1) (CA INDEX NAME) CM 1

RN 512194-04-6 HCAPLUS

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
CN Imidazo(5,1-b:4,3-b']bisoxazol-4-ium, 2,3,7,8-tetrahydro-3,7-bis(phenylmethyl)-, (35,78)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CRN 512194-03-5 CMF C21 H21 N2 O2

Absolute stereochemistry

CM 2

648929-49-1 HCAPLUS Imidazo[5,1-b:4,3-b']bisoxazol-4-ium, 2,3,7,8-tetrahydro-3,7-diphenyl-, (35,78)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-48-0 CMF C19 H17 N2 O2

Absolute stereochemistry

CM 2.

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN Absolute stereochemistry. (Continued)

CM 2

CRN 37181-39-8 CMF C F3 O3 S

648929-57-1 HCAPLUS Imidazo[5,1-b]oxazolium, 2,3-dihydro-3-(1-methylethyl)-6-(2,4,6-trimethylphenyl)-, (3S)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-56-0 CMF C17 H23 N2 O

Absolute stereochemistry.

2 CM

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) CRN 37181-39-8 CMF C F3 O3 S

648929-51-5 HCAPLUS
Imidazo[5,1-b]oxazolium, 2,3-dihydro-3-(1-methylethyl)-6-phenyl-, (3S)-, salt with trifluoromethanesulfonic acid (1:1) [9CI] (CA INDEX NAME)

CRN 648929-50-4 CMF C14 H17 N2 O

Absolute stereochemistry.

648929-53-7 HCAPLUS Imidazo[5,1-b]oxazollum, 3-{1,1-dimethylethyl}-2,3-dihydro-6-phenyl-, (3S)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

648929-59-3 HCAPLUS Imidazo[5,1-b]oxazolium, 3-(1,1-dimethylethyl)-2,3-dihydro-6-(2,4,6-trimethylphenyl)-, (35)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

Absolute stereochemistry.

CM 2

CRN 648929-60-6 CMF C39 H45 N2 O4 Si2

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) CM 2 CRN 37181-39-8 CMF C F3 03 S RN 648929-63-9 HCAPLUS
CN Imidaro[5,1-b:4,3-b']bisoxazol-4-1um,
3,7-bis[diphenyl[(trachylsilyl)oxy]
methyl]-2,3,7,8-tetrahydro-, (3R,7R)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME) CM 1 CRN 648929-62-8 CMF C45 H57 N2 O4 S12 Absolute stereochemistry. CM 2 CRN 37181-39-8 CMF C F3 03 S RN 648929-65-1 HCAPLUS
CN Imidazo[5,1-b]oxazolium,
6-[2,6-bis(1-methylethyl)phenyl]-2,3-dihydro-3-(1methylethyl)-, (3S)-, salt with trifluoromethanesulfonic acid (1:1) [9CI) L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) CM 2 F-C-503 648929-69-5 HCAPLUS Imidazo[5,1-b]oxazolium, 2,3-dihydro-3,6-diphenyl-, (35)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME) CM 1 Absolute stereochemistry. CRN 37181-39-8 CMF C F3 03 S

648929-71-9 HCAPLUS Imidazo[5,1-b]oxazolium, 2,3-dihydro-6-phenyl-3-(phenylmethyl)-, (35)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-70-8 CMF C18 H17 N2 O Absolute stereochemistry. L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (CA INDEX NAME) (Continued) CH 1 CRN 648929-64-0 CMF C20 H29 N2 O Absolute stereochemistry. CM 2 CRN 37181-39-8 CMF C F3 03 S F- c- 503 648929-67-3 HCAPLUS Imidazo[5,1-b]oxazolium, 6-{2,6-bis(1-methylethyl)phenyl}-3-{1,1-dimethylethyl}-2,3-dihydro-, (35)-, salt with trifluoromethanesulfonic acid (1:1) (9C1) (CA INDEX NAME) CM 1 CRN 648929-66-2 CMF C21 H31 N2 O Absolute stereochemistry. L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN 2 648929-73-1 HCAPLUS Imidazo[5,1-b]oxazolium, 2,3-dihydro-3-phenyl-6-(2,4,6-trimethylphenyl)-, (3S)-, salt with trifluoromethanesulfonic acid (1:1) [9CI] (CA INDEX NAME) CM 1 CRN 648929-72-0 CMF C20 H21 N2 O Absolute stereochemistry. CH 2 CRN 37181-39-8 CMF C F3 O3 S

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

648929-75-3 HCAPLUS Imidazo(5,1-b)oxazolium, 2,3-dihydro-3-(phenylmethyl)-6-(2,4,6-trimethylphenyl)-, (35)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-74-2 CMF C21 H23 N2 O

Absolute stereochemistry.

2

CRN 37181-39-8 CMF C F3 O3 S

648929-77-5 HCAPLUS Imidazo[5,1-b]oxazolium, 6-[2,6-bis(1-methylethyl)phenyl]-2,3-dihydro-3-phenyl-, (35)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA IMDEX NAME)

CM 1

CRN 648929-76-4 CMF C23 H27 N2 O

Absolute stereochemistry.

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

CM 2

CRN 37181-39-8 CMF C F3 03 S

648929-79-7 HCAPLUS Imidazo[5,1-b]oxazolium, 6-{2,6-bis(1-methylethyl)phenyl}-2,3-dihydro-3-(phenylmethyl)-, (3S)-, salt with trifluoromethanesulfonic acid {1:1} (9CI) (CA INDEX NAME)

CM 1

CRN 648929-78-6 CMF C24 H29 N2 O

Absolute stereochemistry.

CM 2

CRN 37181-39-8 CMF C F3 O3 S

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

648929-81-1 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-(1,2-ethanediyl)bis[2,3-dihydro-3-{1-methylethyl}-, (35,3's)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

CM 1

Absolute stereochemistry.

2

CRN 37181-39-8 CMF C F3 O3 S

648929-83-3 HCAPLUS
Imidazo[5,1-b]oxazolium, 6,6'-{1,2-ethanediy1}bis[2,3-dihydro-3-(phenylmethyl)-, (3S,3'S)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-82-2 CMF C26 H28 N4 O2

Absolute stereochemistry.

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

2

648929-85-5 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-[1,2-cyclohexanediy1]bis[2,3-dihydro-3-(1-methylethyl)-, (38,3'S)-, salt with trifluoromethanesulfonic acid (1:2) (SCI) (CA INDEX NAME)

CM 1

CRN 648929-84-4 CMF C22 H34 N4 O2

Absolute stereochemistry.

CM 2

CRN 37181-39-8 CMF C F3 O3 S

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L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
                                                          (Continued)
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648929-87-7 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-{1,2-ethanediyl}bis{3-{1,1-dimethylethyl}-2,3-dihydro-, (3S,3'S)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-86-6 CMF C20 H32 N4 O2

Absolute stereochemistry.

CM 2

CRN 37181-39-8 CMF C F3 03 S

648929-89-9 HCAPLUS
Imidazo[5,1-b]oxazolium, 6,6'-(1,2-ethanediyl)bis[2,3-dihydro-3-phenyl-, (35,3'S)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-88-8 CMF C24 H24 N4 O2

ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN CRN 37181-39-8 CMF C F3 03 S (Continued)

648929-93-5 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-[1,2-cyclohexanediyl]bis[2,3-dihydro-3-phenyl-, (38,3's)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

648929-95-7 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-(1,3-phenylene)bis(2,3-dihydro-3-(1-methylethyl)-, (35,3'5)-, selt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-94-6 CMF C22 H28 N4 O2

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN Absolute stereochemistry. (Continued)

CM 2

CRN 37181-39-8 CMF C F3 03 S

RN 648929-91-3 HCAPLUS
CN Imidazo[5,1-b]oxazolium, 6,6'-{1,2-cyclohexanediyl}bis[3-{1,1-dimethylethyl}-2,3-dihydro-, (35,3'5)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-90-2 CMF C24 H38 N4 O2

Absolute stereochemistry

CM 2

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN Absolute stereochemistry. (Continued)

CRN 37181-39-8 CMF C F3 03 S

648929-97-9 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-(1,3-phenylene)bis[2,3-dihydro-3-phenyl-, (35,3'S)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

CM 1

CRN 648929-96-8 CMF C28 H24 N4 O2

Absolute stereochemistry.

2

CRN 37181-39-8 CMF C F3 O3 S

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L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
        648929-99-1 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-[2,6-pyridinediyl]bis[2,3-dihydro-3-[1-methylethyl]-, (35,3'5)-, salt with trifluoromethanesulfonic acid (1:2) (SCI) (CA INDEX NAME)
        CRN 648929-98-0
CMF C21 H27 N5 O2
Absolute stereochemistry
        CRN 37181-39-8
CMF C F3 O3 S
RN 648930-01-2 HCAPLUS
CN Imidazo[5,1-b]oxazolium,
6,6'-(2,6-pyridinediyl)bis[2,3-dihydro-3-phenyl-,
(35,3's)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)
L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
                                                                                                       (Continued)
                                                                Pr-i
        CM 2
        CRN 37181-39-8
CMF C F3 O3 S
        648930-05-6 HCAPLUS Imidazo[5,1-b]Oxazolium, 6,6'-methylenebis[2,3-dihydro-3-phenyl-, (3S,3'S)-, salt with trifluoromethanesulfonic acid (1:2) [9CI] (CA INDEX NAME)
        CM 1
        CRN 648930-04-5
CMF C23 H22 N4 O2
Absolute stereochemistry.
        CM 2
        CRN 37181-39-8
CMF C F3 03 S
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648930-07-8 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-{1,2-cyclohexanediyl}bis[2,3-dihydro-3-(phenylmethyl)-, (35,3'5)-, selt with trifluoromethanesulfonic acid (1:2) {9CI} (CA INDEX NAME)

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L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
                                                                                           (Continued)
       CM 1
       CRN 648930-00-1
CMF C27 H23 N5 O2
Absolute stereochemistry
     648930-03-4 HCAPLUS
Imidazo[5,1-b]oxazolium,
'-methylenebis[2,3-dihydro-3-(1-methylethyl)-,
(38,3'5)-, salt with trifluoromethanesulfonic acid (1:2) {9CI} (CA INDEX NAME)
       CM 1
       CRN 648930-02-3
CMF C17 H26 N4 O2
Absolute stereochemistry.
L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
                                                                                           (Continued)
       CM 1
       CRN 648930-06-7
CMF C30 H34 N4 O2
Absolute stereochemistry.
       CRN 37181-39-8
CMF C F3 O3 S
F- C- 503-
RN 648930-10-3 HCAPLUS
CN Imidazo[5,1-b]oxazolium,
6,6'-(1,3-phenylene)bis[3-(1,1-dimethylethyl)-2,3-
dihydro-, (35,3's)-, salt with trifluoromethanesulfonic acid {1:2} (9CI)
(CA INDEX NAME)
       CM 1
       CRN 648930-09-0
CMF C24 H32 N4 O2
Absolute stereochemistry.
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L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STM CRN 37181-39-8 CMF C F3 03 S (Continued)

648930-12-5 HCAPLUS
Imidazo[5,1-b]oxazolium, 6,6'-(1,3-phenylene)bis[2,3-dihydro-3-(phenylmethyl)-, (35,3'S)-, salt with trifluoromethanesulfonic acid (1:2)
(9C1) (CA INDEX NAME)

CRN 648930-11-4 CMF C30 H28 N4 O2

Absolute stereochemistry.

2

CRN 37181-39-8 CMF C F3 O3 S

RN 648930-14-7 HCAPLUS
CN Imidazo[5,1-b]oxazolium,
6,6'-(2,6-pyridinediyl)bis[3-(1,1-dimethylethyl)2,3-dihydro-, (38,3'5)-, salt with trifluoromethanesulfonic acid (1:2)
(9CI) (CA INDEX NAME)

CM 1

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN CRN 648930-13-6 CMF C23 H31 N5 O2 (Continued)

Absolute stereochemistry.

CM 2

CRN 37181-39-8 CMF C F3 O3 S

648930-16-9 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-{2,6-pyridinediyl}bis[2,3-dihydro-3-(phenylmethyl)-, (38,3'S)-, salt with trifluoromethanesulfonic acid (1:2) {9CI} (CA INDEX NAME)

CM 1

CRN 648930-15-8 CMF C29 H27 N5 O2

Absolute stereochemistry.

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

CM 2

648930-18-1 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-methylenebis[3-{1,1-dimethylethyl}-2,3-dihydro-,(35,3'S)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

СМ 2

CRN 37181-39-8 CMF C F3 O3 S

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

648930-20-5 HCAPLUS Imidazo[5,1-b]oxazolium, 6,6'-methylenebis[2,3-dihydro-3-{phenylmethyl}-, (33,3'S)-, salt with trifluoromethanesulfonic acid (1:2) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

2

CRN 37181-39-8 CMF C F3 03 S

RN 648930-30-7 HCAPLUS
CN Imidazo[5,1-b:4,3-b']bisbenzoxazol-11-ium,
1,2,3,4,4a,6a,7,8,9,10,10a,13adodecahydro-, (4aR,6aR,10aS,13aS)-, salt with trifluoromethanesulfonic
acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 648930-29-4 CMF C15 H21 N2 O2

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

2

CRN 37181-39-8 CMF C F3 O3 S

648930-34-1 HCAPLUS Imidazo[5,1-b]oxazolium, 6-[1-[{4S}-4,5-dihydro-4-(1-methylethyl)-2-oxazolyl]-1-methylethyl]-2,3-dihydro-3-(1-methylethyl)-, (3S)-, salt with trifluoromethanesulfonic acid (1:1) [9CI] (CA INDEX NAME)

CH 1

CRN 648930-33-0 CMF C17 H28 N3 O2

Absolute stereochemistry

2

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

L69 ANSWER 10 OF 63
ACCESSION NUMBER:
DOCUMENT NUMBER:
139:276684
An N-heterocyclic carbene ligand with flexible steric bulk allows Suzuki cross-coupling of sterically hindered aryl chlorides at room temperature
Altenhoff, Gereenn Goddard, Richard; Lehmann, Christian W. Glorius, Frank
MAX-Planck-Institut fuer Kohlenforschung, Muelheim an der Ruhr, 45470, Germany
SOURCE:

PUBLISHER:
PUBLISHER:
PUBLISHER:
BUILISHER:
PUBLISHER:
PUBLISHER:
BUILISHER:
ANGURGE:
AN

August 11,2003

OTf

A catalyst prepared from Pd(OAc)2 and imidazolium salt I catalyzed the Suzuki cross-coupling of sterically hindered and unhindered, activated

unactivated, aryl chlorides and aryl boronic acids. Obtained were di-

Tri-ortho-substituted biphenyl compds.

606970-69-8P
RL: CAT (Catalyst use): PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (crystal structure; Suruki cross-coupling of sterically hindered aryl chlorides and aryl boronic acids catalyzed by catalyst prepared from Pd(OAc)2 and imidazolium selt)

606970-69-8 HCAPLUS
Dispiro(cyclohexane-1,2'(2'H)-imidazo[5,1-b:4,3-b']bisoxazol[4]ium-7'(8'H),1''-cyclohexane}, selt with trifluoromethanesulfonic acid (1:1)

(9CI) (CA INDEX NAME)

CH 1

CRN 606970-68-7 CMF C17 H25 N2 O2

L69 ANSWER 9 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT: THIS

THERE ARE 10 CITED REFERENCES AVAILABLE FOR

FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L69 ANSWER 10 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 34 CITED REFERENCES AVAILABLE FOR

FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L69 ANSWER 11 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER: 2003:563667 HCAPLUS
DOCUMENT NUMBER: 140:76926
TITLE: CP0569. A New Record Control of the Control o

CP0569, A New Broad-Spectrum Injectable Carbapenem. Part 1: Synthesis and Structure-Activity

CORPORATE SOURCE:

Relationships AUTHOR(S):

Aihara, Kazuhiro; Kano, Yuko; Shiokawa, Sohjiro; Sasaki, Toshiro; Setsu, Fumihito; Sambongi, Yumiko; Ishii, Miyuki; Tohyama, Kazuyo; Ida, Takashi; Tamura, Atsushi; Atsumi, Kunio; Iwamatsu, Katsuyoshi Pharmaceutical Research Center, Heiji Selka Kaisha, Ltd., Kohoku-ku, Yokohama, 222-8567, Japan Bioorganic & Medicinal Chemistry (2003), 11(16), 3475-3485

SOURCE: August 5

CODEN: BMECEP; ISSN: 0968-0896 Elsevier Science Ltd.

PUBLISHER:

DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 140:76926

AB A series of 1B-methylcarbepenems bearing an (imidazo[5,1-b]thiazolium6-yllmethyl moiety, a 5,5-fused heterobicycle, at the C-2 position was
synthesized and evaluated for in vitro antibacterial activities. CPD569
(Ir) and its analogs showed potent antibacterial activities against
Gram-pos bacteria, including methicillin-resistant Staphylooccus aureus
(MRSA), and Gram-neg. bacteria, including Pseudomonas aeruginosa.
Moreover, CPD569 (Ir) exhibited stronger antibacterial activity against
MRSA and higher resistance to renal dehydropeptidase-1 (DHP-1) than any
currently marketed carbapenems, i.e., imipenem (IPM), panipenem (PAPM),
and meropenem (MEPM).

IT 640275-17-89 640275-19-0P
RL: PRAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)

(Uses)

(synthesis and structure-activity relationships of carbapenem CP0569)

RN 640275-17-8 HCAPLUS

(Indiazol5,1-b]thiazolium,
6-{(46,5,8,65)-2-carboxy-6-[(1R)-1-hydroxyethyl)4-methyl-7-oxo-1-azabicyclo{3.2.0}hept-2-en-3-yl}methyl]-2,3-dihydro-,
inner salt (9C1) (CA INDEX NAME)

Absolute stereochemistry.

640275-19-0 HCAPLUS Imidazo[5,1-b]benzothiazolium, 2-[[(48,5R,68)-2-cerboxy-6-[(1R)-1-hydroxyethyl]-4-methyl-7-oxo-1-azabicyclo[3.2.0]hept-2-en-3-yl]methyl-, inner salt (9CI) (CA INDEX NAME)

L69 ANSWER 11 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) Absolute stereochemistry.

REFERENCE COUNT: THIS

FORMAT

14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

L69 ANSWER 12 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
138:321365
TITLE:
CNAROLINES as chiral building blocks for imidazolium salts and N-heterocyclic carbene ligands
AUTHOR(S):
CORPORATE SOURCE:
MAX-Planck-Institut fuer Kohlenforschung,
Nuelheim/Ruhr, 45470, Germany
CORDICE:
Chemical Communications (Cambridge, United Kingdom)
(2002), (22), 2704-2705
CODEN: CHCOPS; ISSN: 1359-7345
PUBLISHER:
BOUNDERT TYPE:
DOCUMENT TYP RE: CAT (Catalyst use); PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (preparation and crystal structure of enantiomerically pure oxazoline-based oline-based imidazolium triflates and their deprotonation to chiral N-heterocyclic carbenes as ligands for palladium-catalyzed arylation reactions) 512193-98-5 HCAPLUS IMidazo[5,1-b:4,3-b']bisoxazol-4-ium, 2,3,7,8-tetrahydro-3,7-bis(1-methylethyl)-, (35,75)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME) CM 1

CRN 512193-97-4 CMF C13 H21 N2 O2 Absolute stereochemistry. Rotation (+).

CM 2 CRN 37181-39-8 CMF C F3 03 S

512194-01-3P 512194-04-6P RL: CAT (Catalyst use); SPN (Synthetic preparation); PREP (Preparation); KL: CAT (Catalyst use); SPN (Synthetic preparation); PREF (Preparation USES (Uses)
(preparation of enantiomerically pure oxazoline-based imidazolium triflates lates
and their deprotonation to chiral N-heterocyclic carbenes as ligands
for palladium-catalyzed arylation reactions)
512194-01-3 HCAPLUS
Imidazo[5,1-b:4,3-b\*]bisoxazol-4-ium, 3,7-bis(1,1-dimethylethyl)-2,3,7,8tetrahydro-, (3S,7S)-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME) CM 1 CRN 512194-00-2 CMF C15 H25 N2 O2 Absolute stereochemistry CM 2 CRN 37181-39-8 CMF C F3 03 S F- C- SO3-512194-04-6 HCAPLUS Imidazo[5,1-b:4,3-b']bisoxazol-4-ium, 2,3,7,8-tetrahydro-3,7-bis(phenylmethyl)-, (35,78)-, salt with trifluoromethanesulfonic acid (1:1) (9C1) (CA INDEX NAME) CH 1 CRN 512194-03-5 CMF C21 H21 N2 O2 Absolute stereochemistry.

L69 ANSWER 12 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

L69 ANSWER 12 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

CM 2

CRN 37181-39-8 CMF C F3 03 S

IT 512194-12-6P 512194-15-9P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of enantiomerically pure oxazoline-based imidazolium triflates

lates
and their deprotonation to chiral N-heterocyclic carbenes as ligands
for palladium-catalyzed arylation reactions)
512194-12-6 HCAPEUS
2H-Imidazo(5,1-b]oxazo1-4-ium, 3,6-dihydro-3-(1-methylethyl)-6-(2,4,6trimethylphenyl)-, (3S)-, salt with trifluoromethanesulfonic acid (1:1)
(9CI) (CA INDEX NAME)

CRN 512194-11-5 CMF C17 H23 N2 O

Absolute stereochemistry

CM 2

L69 ANSWER 12 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN RN 512194-17-1 HCAPLUS CN 5H-Imidazo[5,1-b:4,3-b']bisoxazol-5-ylidene, 2,3,7,8-tertaphydro-3,7-bis(1-methylethyl)-, (3S,7S)- (9CI) (CA INDEX NAME) (Continued)

Absolute stereochemistry.

REFERENCE COUNT: THIS

22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 12 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN CRN 37181-39-8 CMF C F3 03 S (Continued)

512194-15-9 HCAPLUS 2H-Imidazo[5,1-b]oxazol-4-ium, 6-[2,6-bis(1-methylethyl)phenyl]-3,6-dihydro-3-(1-methylethyl)-, {3S}-, salt with trifluoromethanesulfonic

(1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 512194-14-8 CMF C20 H29 N2 O

Absolute stereochemistry.

2

CRN 37181-39-8 CMF C F3 03 S

IT 512194-17-1P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or respent)
(reaction with sulfur; preparation of enantiomerically pure oxazoline-based indazolium triflates and their deprotonation to chiral N-heterocyclic carbenes as ligands for palladium-catalyzed arylation reactions)

L69 ANSWER 13 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN

ACCESSION NUMBER: 2002;379196 HCAPLUS

TITLE: An expedient method for the solid-phase synthesis of α-aminoalkyl phosphonopeptides

AUTHOR(S): Rinnova, Marketa; Nefzi, Adel; Houghten, Richard A. Torrey Pines Institute for Molecular Studies, San Diego, CA, 92121, USA

SOURCE: TELEAY; ISSN: 0040-4039

PUBLISHER: DESevier Science Ltd.

DOCUMENT TIPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 137:201560

AB The formation of α-amino phosphonate functionalities on the amino terminus of peptides utilizing solid-phase methodol. is presented. The described method enables incorporation of diverse N-phosphonoalkyl and aryl moleties.

I 45340-91-5P

RL: SPN (Synthetic preparation); PREF (Preparation)

(solid-phase synthesis of aminoalkyl phosphonopeptides)

RN 453540-91-5 HCAPLUS

Absolute stereochemistry.

THERE ARE 29 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

REFERENCE COUNT:

L69 ANSWER 14 OF 63
ACCESSION NUMBER:
DOCUMENT NUMBER:
137:6120
Highly diastereoselective addition of
N-Boc-pyrrolidin-2-yllithium to optically active
ketimines - synthesis of enantiometically pure
1,3-imidarolidin-2-ones and diamines
Von Keyserlingk, Nikolai Graf; Martens, Jurgen
Universitat Oldenburg, Fachbereich Chemie, Oldenburg,
26129, Germany
SOURCE:

301-308 CODEN: EJOCFK; ISSN: 1434-193X Wiley-VCH Verlag GmbH Journal English CASREACT 137:6120 FUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
GI

PUBLISHER:

AB A highly disstereoselective addition of chiral

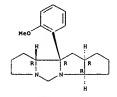
N-Boc-pytrolidin-2-yllithium
to optically active bicyclic ketimines has been developed. For this
purpose slkyl- and aryl-substituted chiral N-Boc-amino ketones have been
synthesized by addition of various Grignard reagents to an

N-Boc-protected
lactam. The resulting N-Boc-amino ketones have been converted into
bicyclic ketimines after deprotection and intramol. cyclization. A
kinetic resolution of the racemic organolithium compound by the chiral
substrate is discussed based on x-ray crystal structure anal. and exptl.
results. The influence of the substituent of the ketimine has been
studied. Some of the obtained tetracyclic 1,3-imidatolidin-2-ones I (R =
Ph, 4-McC6H4, 3-McC6H4, 2-McC6H4).

IT 431887-13-7P 431887-41-1P 431887-43-3P
RL: RCT (Reactant): SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(diastereoselective addition of N-Boc-pyrrolidin-2-yllithium to
optically
active ketimines for synthesis of enantiomerically pure
1,3-imidazolidin-2-ones and diamines and kinetic resolution of the
racemic

organolithium compound) 431887-13-7 HCAPLUS

L69 ANSWER 14 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN



431887-42-2P
RL: SPN (Synthetic preparation); PREP (Preparation)
(diastereoselective addition of N-Boc-pyrrolidin-2-yllithium to

optically
active ketimines for synthesis of enantiomerically pure
1,3-imidazolidin-2-ones and diamines and kinetic resolution of the nic organolithium compound)
431887-42-2 HCAPLUS
1H, 5H-Cyclopenta[4,5]pyrrolo[1,2-c]pyrrolo[2,1-e]imidazole,
decahydro-9b-(3-methoxyphenyl)-, (3aR,9aR,9bR,10aR)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

20

REFERENCE COUNT:

THERE ARE 20 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

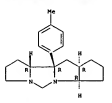
ANSWER 14 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (ContinuitH,5H-Cyclopenta(4,5)pyrrolo[1,2-c]pyrrolo[2,1-c]imidazole, decahydro-9b-phenyl-, (3a, 9a, 9a, 10aR)- (9CI) (CA INDEX RAME) (Continued)

Absolute stereochemistry.

431887-41-1 HCAPLUS

1H, 5H-Cyclopenta[4,5]pyrrolo[1,2-c]pyrrolo[2,1-e]imidazole, decahydro-9b-{4-methylphenyl}-, (3aR,9aR,9bR,10aR)- (9CI) (CA INDEX NAME )

Absolute stereochemistry. Rotation (+).



431887-43-3 HCAPLUS

HILDER TO THE STATE OF T

Absolute stereochemistry. Rotation (+).



L69 ANSWER 15 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2001:905331 HCAPLUS
DOCUMENT NUMBER: 136:241071
TITLE: Increased rigidity of the chiral centre of tocainide favours stereoselectivity and use-dependent block of skeletal muscle Na+ channels enhancing the antimyotonic activity in vivo
AUTHOR(S): Talon, Sophie; De Luca, Annamaria; De Bellia,

AUTHOR(S): Michela;

Desaphy, Jean-Francois: Lentini, Giovanni; Scilimati, Antonio: Corbo, Filomena: Franchini, Carlo; Tortorella, Paolo; Jockusch, Harald; Camerino, Diana

Tortorelia, Paolo; Jockusch, Harald; Camerino, Diana Conte

CORPORATE SOURCE: Department of Pharmacobiology, Unit of Pharmacology, Faculty of Pharmacy, University of Bari, Bari, 1-70125, Italy

SOURCE: British Journal of Pharmacology (2001), 134(7), 1523-1531

CODDN: BJPCBN; ISSN: 0007-1188

PUBLISHER: Nature Publishing Group
Journal
LANGUAGE: English

AB 1 Searching for the structural requirements improving the potency and the stereoselectivity of Nar channel blockers as antimyotonic agents, new derivs. of tocainide, in which the chiral carbon atom is constrained in a rigid a-proline or pyrrolo-imidazolic cycle, were tested as pure enantiomers. 2 Their ability to block Na+ currents, elicited from -100 to

enantiomers. 2 Their ability to block Na+ currents, elicited from -100 to -20 mV at 0.3 Hz (tonic block) and 2-10 Hz (use-dependent block) frequencies, was investigated in vitro on single fibers of frog semitendinosus muscle using the vaseline-gap voltage-clamp method. 3 The α-proline derivative, 765, was 5 and 21 fold more potent than tocainide in producing tonic and 10 Hz-use-dependent block, resp. Compared to 765, the presence of one Ne group on the aminic (766) or amidic (767) nitrogen atom decreased use-dependence by 2- and 6-times, resp. When methylene moieties were present on both nitrogen atoms (768), both tonic and use-dependent block were reduced. 4 Contrarily to tocainide, all proline derivs. were stereoselective in relation to an increased rigidity, A further increase in the mol. rigidity as in pyrrolo-imidazolic derivs. markedly decreased the drug potency with respect to tocainide. 5 Antimyotonic activity, evaluated as the shortening of the time of righting rellexes of myotonic adr/adr mice upon acute drug in vivo administration was 3 fold more effective for R-To5 than for R-Tocainide. 6 Thus, constraining the chiral center of tocainide in α-proline cycle leads to more potent and stereoselective use-dependent Nav channel blockers with

improved therapeutic potential.
403995-20-0 403995-21-1
RL: PAC (Pharmacological activity): THU (Therapeutic use): BIOL
(Biological study): USES (Uses)
(increased rigidity of the chiral center of tocainide favors
stereoselectivity and use-dependent block of skeletal muscle Na+
channels enhancing the antimyotonic activity in vivo)
403995-20-0 HCAPLUS
HI-Pyrrolofi, 2-climidazol-1-one, 2-(2,6-dimethylphenyl)hexahydro-, (7aR)(9CI) (CA INDEX NAME)

L69 ANSWER 15 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

403995-21-1 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, 2-(4-chloro-2-methylphenyl)hexahydro-,
(7aR)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

THERE ARE 33 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

ANSWER 16 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
Carbapenem derivs. represented by the general formula [I; Rl = H, Me; R2, R3 = H, halo, lower alkyl optionally substituted by HO or NHZ, lower alkylcathonyl, CONHZ, aryl, lower alkylcathoic, R4 = (un)substituted lower alkylthio, lower cycloalkylthio, C2-4 alkenylthio, C2-4 alkynylthio,

alkylthio, lower cycloalkylthio, C2-4 alkenylthio, C2-4 alkynylthio, monoor bicyclic heterocyclylthio containing 21 of same or different heteroatoms, lower alkylsulfinyl, (un)substituted lower alkylsulfonyl, lower alkylcarbonyl, arylcarbonyl; or R4 and R5 are linked to each other to represent S(CR2)n (n = 2-4); R5 = (un)substituted lower alkyl, lower cycloalkyl, C2-4 alkenyl, C2-4 alkynyl, (un)substituted 4- to 7-membered aliphatic heterocyclyl optionally containing 21 of 0 or S atoms] are prepared These compds. have potent antibacterial activities on methicillin-resistant Staphylococcus aureus (MRSA), penicillin-resistant Streptococcus pneumoniae (PRSP), Haemophilus influenzae, and β-lactamase-producing bacteria and a high stability to renal dehydropeptidase enzyme (DHP-1). Thus, (15,5M,65)-6-f(1R1-1-hydroxyethyl)1-methyl-2-(7-methylthioimidazo(5,1-b)thiazol-2-yl)-1-carbapen-2-em-3-carbonylic acid p-nitrobenzyl ester (preparation given) was dissolved in CH2C12, cooled in an ice bath, treated with 0.022 mL Me trifluoromethanesulfonate, and stirred at the same temperature for 30 min to

min

to give (18,5R,6S)-6-[(1R)-1-hydroxyethyl]-1-methyl-2-(6-methyl-7-methylthioimidazo(5,1-b]thiazolium-2-yl)-1-carbapen-2-em-3-carboxylic

acid

p-nitrobenzyl ester trifluoromethanesulfonate which was hydrogenolyzed

over 10% Pd-C in a mixture of 1 N phosphate buffer (pH 6.8) and THF under
hydrogen atmospheric for 1.5 h to give
(15.5%,65)-6-[(1%)-1-hydroxysthyl]-1-

methyl-2-(6-methyl-1-methylthioimidazo[5,1-b]thiazolium-2-yl)-1-carbapen-2-em-3-carboxylate (inner salt) (II). II in vitro showed min. inhibitory concentration of 1.56 and 0.025 µg/mL against highly methicillin-resistant
Staphylecoccus aureus M126 and highly penicillin-resistant Streptococcus pneumoniae, resp.

1 35306-76-4P
RL: BAC (Biological activity or effector, except adverse); BSU

(Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of novel carbapenem derivs. quaternary salts as antimicrobial

nicrobial
agents)
352306-76-4 HCAPLUS
352306-76-4 HCAPLUS
Imidazo[5,1-b:4,3-b']bisthiazol-4-ium, 8-[(45,5R,6S)-2-carboxy-6-[(1R)-1-hydroxyethy]1-4-methyl-7-oxo-1-azabicyclo[3,2.0]hept-2-en-3-yl]-2,3dihydro-, inner salt (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 16 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 2001:565047 HCAPLUS DOCUMENT NUMBER: 135:152661

TITLE:

135:152661
Preparation of novel carbapenem derivatives of quaternary salt type as antimicrobial agents Kano, Yuko: Maruyama, Takahisa; Yamamoto, Yasuo; Shitara, Eiji; Sasaki, Toshiro; Aihara, Kazuhiro; Atsumi, Kunio; Iwamatsu, Katsuyoshi; Ida, Takashi Mekji Seika Kaisha, Ltd., Japan PCT Int. Appl., 329 pp. CODEN: PIXXD2
Patent INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA'	TENT :	NO.			KIN	D	DATE									ATE	
						-									-		
WO	2001	0551	55		A1		2001	0802		WO 2	001-	JP52	9		2	0010	126
	W:																
		CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
		ΗU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,
		LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
		SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	υs,	UZ,	VN,
		YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM				
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	G₩,	ML,	MR,	NE,	SN,	TD,	TG		
ΕP	1251	134			A1		2002	1023		EP 2	001-	9468	65		2	0010	126
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		IE,	SI,	LT.	LV,	FI,	RO,	MK,	CY,	AL,	TR						
										US 2	002-	1821	80		2	0020	725
US	6825	187			B2		2004	1130									
RIT	Y APP	LN.	INFO	. :						JP 2	000-	1741	8		A 2	0000	126
										WO 2	001-	JP52	9	,	w 2	0010	126
	CA AU EP US	CA 2398 AU 2001 EP 1251 R: US 2003 US 6825	W0 20010551 W: AE, CR, HU, LU, SD, YU, RW: GH, BJ, CA 2398478 AU 20010288 EP 1251134 R: AT, IS, US 20030228	WO 2001055155 W: AE, AG,	W: AE, AG, AL, CR. CU, CZ, HU, ID, IL, LU, LV, MA, SD, SE, SG, YU, ZA, ZW, RW: GH, GM, KE, DE, DK, ES, BJ, CF, CG, CA 2398478 R: AT, BE, CH, IE, SI, LT, US 2003022881	M: AE, AG, AL, AM, CR, CU, CZ, DE, HU, ID, IL, IN, LU, LV, MA, MG, SD, SE, SG, MG, YU, ZA, ZW, AM, RW: GM, GM, KE, FI, BJ, CF, CG, CI, CA 2398478 AA AU 2001028833 AB FF 1251134 A1 R: AT, BE, CH, DE, 1E, SI, LT, LV, US 2003022891 A1 US 6825187 B2	M: AE, AG, AL, AM, AT, CR. CU, CZ, DE, DK, HU, ID, IL, IN, IS, LU, LV, MA, MD, MG, SD, SE, SG, ST, SK, YU, ZA, ZW, AM, AZ, RW: GM, GM, KE, LS, MM, DE, DK, ES, FI, FR, BJ, CF, CG, CI, CM, CA 2398478 AA AU 2001026833 A5 EP 1251134 A1 R: AT, BE, CH, DE, DK, IE, SI, LT, FI, US 2003022891 A1 US 6825187 B2	WC 2001055155 A1 2001 W: AE, AG, AL, AM, AT, AU, CR, CU, CZ, DE, DK, DM, HU, IO, IL, IN, IS, JP, LU, LV, MA, MD, MG, MK, SD, SE, SG, SI, SK, SL, YU, ZA, ZW, AM, AZ, BY, DE, DK, ES, FI, FR, GB, BJ, CF, CG, CI, CM, AD, CA 2398478 AA 2001 AU 2001028833 A5 2001 EP 1251134 A1 2002 R: AT, BE, CH, DE, DK, ES, IE, SI, IV, FIR, GS, CS, CS, CS, CS, CS, CS, CS, CS, CS, C	W: AE, AG, AL, AM, AT, AU, AZ, W: AE, AG, AL, AM, AT, AU, AZ, C, CU, CZ, DE, DK, DM, DZ, HU, ID, IL, IN, IS, JP, KE, LU, LV, MA, MD, MG, MK, MN, SD, SE, SG, SI, SK, SL, TJ, YU, ZA, ZW, AM, AZ, BY, KG, DE, DK, ES, FI, FR, GB, GR, CA 2398478 AA 20010807 EP 1251134 A1 20021023 R: AT, BE, CH, DE, DK, ES, FR, IE, SI, LT, LV, FI, RO, MK, US 2003022881 A1 20030313	WC 2001055155 A1 20010802  W: AE, AG, AL, AM, AT, AU, AZ, BA, CR, CU, CZ, DE, DK, DM, DZ, EE, HU, ID, II, IN, IS, JP, KE, KG, LU, LV, MA, MD, MG, MK, MN, MN, SD, SE, SG, SI, SK, SL, TJ, TM, YU, ZA, ZW, AM, AZ, BY, KG, KZ, RW: GH, GM, KE, LS, MM, MZ, SD, SL, DE, DK, ES, FI, FR, GB, GR, IE, BJ, CF, CG, CI, CM, GA, CM, GM, CA 2398478 AA 20010807  EP 1251134 A1 20021023  R: AT, BE, CH, DE, DK, ES, FR, GB, IE, SI, LT, LV, FI, RO, MK, CT, US 2003022881 A1 20030130  USG 6252187 B2 20041130  RITY APPLN. INFO::	WO 2001055155 A1 20010802 WO 2  W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, HU, ID, II, IN, IS, JP, KE, KG, KP, LU, LV, MA, MD, MG, MK, NN, MM, MX, SD, SE, SG, SI, SK, SL, TJ, TM, KY, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, DE, DK, ES, FI, FR, GB, GR, IE, IT, BJ, CF, CG, CI, CM, GA, GN, GW, ML, CA 2398478 AA 20010802 CA 2  AU 2001028833 A5 20010807 AU 2  EP 1251134 A1 20021023 EP 2  ER AT, BE, CH, DE, DK, ES, FR, GB, GR, CM, LE, LT, LY, FI, RO, MK, CY, AL, US 2003022881 A1 20030130 US 2  US 6825187 B2 20041130 F2  RITY APPLN. INFO: : JP 2	WO 2001055155 A1 20010802 WO 2001—  W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, DE, DK, ES, FI, FR, GB, GR, IE, LT, LU, BJ, CF, CG, CT, CM, GA, GN, GW, LM, MR, CA 2398478 AA 20010802 WILL, MR, KG, AZ, DG, CA 2001—  R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, TE, LT, LV, FI, RO, MK, CY, AL, TR US 200301281 A1 20030130 US 2002—  RITY APPLN. INFO:: JP 2000—	MO 2001055155 A1 20010802 MO 2001-JP52  W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FIT, GB, HU, ID, IL, IM, IS, JP, KE, KG, KP, KR, KZ, LU, LV, MA, MD, MG, MK, NN, MM, MX, MZ, NO, SD, SE, SG, SI, SK, SL, TJ, TM, TT, TZ, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, RW: GB, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, DE, DK, ES, FIT, FR, GB, GR, IE, ITT, LU, MC, BJ, CF, CG, CT, CM, GA, GM, GM, LM, KR, NS, CA 2398478 A2 20010807 A2 2001-2838 A2 20010807 A2 2001-2838 A2 20010807 A2 2001-2838 A2 20021023 EP 2001-9468 A2 2003022881 A1 20030130 US 2002-1821 US 6825187 B2 20041130 US 2002-1821 RITY APPLN. INFO::	W0 2001055155 A1 20010802 W0 2001—JF529 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, HU, ID, II, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LU, LV, MA, MD, MG, MK, NN, MW, MX, MZ, ND, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TG, CM, CB, CB, CB, CB, CB, CB, CB, CB, CB, CB	WO 2001055155 A1 20010802 WO 2001-JP529  W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FT, GB, GD, GE, HU, ID, II, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LU, LV, MA, MD, MG, MK, MN, MM, MX, MZ, ND, CY, PL, SD, SE, SG, SI, SK, SL, TJ, TH, TT, TT, TZ, UA, UG, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, ML, PT, BJ, CF, CG, CI, CM, GA, CM, GM, ML, KR, NE, SN, TD, CA 2398478 AA 20010802 CA 2001-2398478  EP 1251134 A1 20021023 EP 2001-946865  EP 1251134 A1 20021023 EP 2001-946865  ER: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, US 2003022881 A1 20030130 US 62625187 B2 20041130 RITY APPLN. INFO::	WO 2001055155  A1 20010802 WO 2001-JP529 2  W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, CH, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LU, LV, MA, MD, MG, MK, NN, MM, MX, NO, NZ, PL, PT, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, RW: GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, BJ, CF, CG, CI, CM, GA, GN, GM, ML, MR, NS, NT, TD, TG CA 2398478  AA 20010802 CA 2001-2398478  AB 20010807  AB 20010807  AB 2001080833  AS 20010807  AS 20010807  AB 20010807  AB 2001080833  BY 1251134  AI 20021023  BY 2001-946865  AB 2003022881  AI 20030130  AI 20030128813  AI 20030130  AI 20030130  AI 2003012810  AI 20030130  AI 2003012810  AI 20030130  AI 20030130  AI 20030130  AI 20030130  AI 2000-17418  AI 2	W0 2001055155

OTHER SOURCE(S): MARPAT 135:152661

L69 ANSWER 16 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

L69 ANSWER 17 OF 63
ACCESSION NUMBER:
DOCUMENT NUMBER:
134:280528
Enantioselective desymmetrization of meso-cyclic anhydrides catalyzed by hexahydro-1H-pyrrolo[1,2-c]imidazolones
AUTHOR(S):
CORPORATE SOURCE:
Uorumi, Y.; Yasoshima, K.; Miyachi, T.; Nagai, S.-i.
Institute for Molecular Science, Myodaiji, Okazaki,
444-8585, Japan
Etrahedron Letters (2001), 42(3), 411-414
CODDE: TELERY: ISSN: 0040-4039
Elsevier Science Ltd.
Journal

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): AB Asym. methan ISBER: Elsevier Science Ltd.

WENT TYPE: Journal

UAGE: English

R SOURCE(S): CASREACT 134:280528

Asym. methanolysis of meso cyclic carboxylic anhydrides including
hexahydrophthalic anhydride proceeded in toluene in the presence of

(6R,7aS)-2-aryl-6-hydroxyhexahydro-1H-pyrrolo[1,2-c]imidazol-1-one to

the corresponding desymmetrized monoester acids, e.g. [15,2R]-2-(methoxycarbonyl)cyclohexane-1-carboxylic acid, with £898 ee.

173549-74-1P 332123-98-5P 332123-99-6P
33293-68-1P
RL: CAT (Catalyst use); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)
(asym. methanolysis of meso-cyclic anhydrides catalyzed by hexahydropyrrolo[1,2-c]imidazolones)
173549-74-1 HCAPJUS
1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-6-hydroxy-2-phenyl-, (6R,7aS)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

332123-98-5 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-phenyl-, (7as)- (9CI) (CA INDEX NAME)

Absolute stereochemistry

332123-99-6 HCAPLUS

L69 ANSWER 17 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN CN 1H-Pyrrolo[1,2-c]imidazol-1-one, 6-[{[1,1-dimethylethyl)dimethyleilyloxy] hexahydro-2-phenyl-, (6R,7aS)- (9CI) (CA INDEX NAME

Absolute stereochemistry.

332943-88-1 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-6-hydroxy-2-(4-octylphenyl)-, (6R, 7as)- (9C1) (CA INDEX NAME)

Absolute stereochemistry.

FORMAT

THERE ARE 36 CITED REFERENCES AVAILABLE FOR

06/28/2006

(Continued)

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L69 ANSWER 18 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2001:47297 HCAPLUS DOCUMENT NUMBER: 134:266218
TITLE: A parallel preparation of a bicy

134:266218

A parallel preparation of a bicyclic N-chiral amine library and its use for chiral catalyst screening Uozumi, Y., Mizutani, K.; Nagai, S.-i.

Institute for Molecular Science, Myodaiji, Okazaki, 444-8585, Japan

Tetrahedron Letters (2001), 42(3), 407-410 CODEN: TELEAY; ISSN: 0040-4039 Elsevier Science Ltd. AUTHOR(S): CORPORATE SOURCE:

SOURCE:

PUBLISHER:

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): English CASREACT 134:266218

A parallel library of optically active bicyclic tertiary amines bearing N-chiral bridgehead nitrogen atoms was readily prepared by condensation

primary amines, cyclic amino acids, and aldehydes. The

enanticontrolling ability of each of the library members was examined for the asym. alkylation

lation
of benzaldehyde with diethylrinc, and (3R,6R,7aS)-(2,3-diphenyl-6-hydroxylhexahydro-lH-pyrrolo(1,2-c|midazol-l-one, which contains a fl-amino alot unit, showed high enantioselectivity.
173549-74-1P 332123-92-9P 332123-95-SP
332123-96-8P 332123-97-4P 332123-98-SP
332123-99-6P

RL: CAT (Catalyst use); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

USES (Uses) (parallel preparation of bicyclic N-chiral amine library) 173549-74-1 HCAPLUS H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-6-hydroxy-2-phenyl-, (6R,7aS)-(9CI) (CA INDEX NAME)

Absolute stereochemistry

332123-92-9 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-6-hydroxy-2-propyl-, (6R,7aS)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 332123-95-2 HCAPLUS
CN 1H-Pyrrolo[1,2-c]imidazol-1-one,
2-(2,6-dimethylphenyl)hexahydro-6-hydroxy-

L69 ANSWER 18 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN , (6R,7as)- (9CI) (CA INDEX NAME) (Continued)

Absolute stereochemistry.

RN 332123-96-3 HCAPLUS CN 1H-Pyrrolo[1, 2-c]imidazol-1-one, 2-(3,5-dimethylphenyl)hexahydro-6-hydroxy-, (6R,7aS)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

332123-97-4 HCAPLUS
HH-Pyrrolo[1,2-e]imidazol-1-one, hexahydro-6-hydroxy-2-(1-naphthaleny1)-, (68,7as)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

332123-98-5 HCAPLUS
1H-Pyrrolo[1, 2-c|imidazol-l-one, hexahydro-2-phenyl-, (7a8)- (9CI) (CA
INDEX NAME)

L69 ANSWER 18 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 332123-99-6 HCAPLUS
CN HR-Pyrrolo[1,2-c]imidazol-1-one,
6-[{[1,1-dimethylethyl]dimethylsilyl]oxy|
hexahydro-2-phenyl-, (6R.7aS)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 19 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

what are claimed are pyrimidine compds. (shown as I), or their pharmaceutically acceptable salts, hydrates, solvates, crystal forms and individual diastereomers, and pharmaceutical compns. including the same and their use as inhibitors of tyrosine kinase enzymes and consequently their use in the prophylaxis and treatment of protein tyrosine kinase-associated disorders, such as immune diseases, hyperpoliferative disorders and other diseases in which inappropriate protein kinase action is believed to play a role, such as cancer, angiogenesis, erosclerosis, graft rejection, rheumatoid arthritis and psoriasis. In I, R1, R2 = independently H, halo, OH, SH, CN, NO2, alkyl, alkoxy, acyloxy, alkoxycathonyloxy, carbamyolxy, alkythio, sulfinyl, sulfonyl, acyl, alkoxycathonyl, carbamoyl, amino, acylamino, ureido, sulfamoyl, sulfonylamino, or R1 and R2 can join together to form a fused methylenedioxy ring or a fused 6-membered aromatic ring; terms such as 'alkyl' here and below are further defined in the claims. R3, R5 = independently H, C1-C6-alkyl unsubstituted or substituted with 1-3 substituents, aryl, or R3 and R5 taken together can represent :0. R3 or

can represent a 2 or 3 C methylene bridge forming a ring of 5-8 atoms fused to the A ring. R4 = H, Cl-C6-alkyl, Cl-C6-alkoxyl. X1, X2, X3, X4 in -X1:X2-X3:X4- are substituted or unsubstituted CH or N where 0-2 of

X1, X2, X3, X4 are N. X5, X6 = independently N, C, optionally substituted

CH. A ring = Ph, naphthyl, pyridyl, pyrazinyl, pyrimidinyl, pyrrolyl,

A ring = Ph, naphthyl, pyridyl, pyrazlnyl, pyrimidinyl, pyrrolyl, thienyl, oxazolyl, isoxazolyl, thiazolyl, pyrazolyl, triazolyl, tetrazolyl, furanyl, benzothienyl, benzofuranyl, indolyl, imidazolyl, benzimidazolyl, thiadiazolyl, R7, R8, R9, R10 = independently H, halo, OH, SH, CN, NO2, N3, N2+BF4-, alkyl, alkoxy, alkylthio, sulfinyl, sulfonyl, C1-C6-perfluoroalkyl, acyl, alkoxycarbonyl, carbamoyl, acyloxy, alkoxycarbonyloxy, carbamoyloxy, and koxycarbonyloxy, carbamoyloxy, amino, acylamino, ureido, sulfamoyl, sulfonylamino, two of R7, R8, R9, and R10 when on adjacent carbons join together to form a methylenedioxy bridge. N = O-2. More than 500 example

ple
prepns. are given, but no preparative method is claimed and no data
relating to the usefulness of the compds. are given.
317826-43-0P, 2-[(S)-(3-Trifluoromethylphenyl)ethylamino]-4-[5-(3-

L69 ANSWER 19 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2001:12273 HCAPLUS

134:86271

DOCUMENT NUMBER: TITLE: 144:862/1 Preparation of pyrimidine derivatives as Src-family protein tyrosine kinase inhibitor compounds Armstrong, Helen M.; Beresia, Richard; Goulet, Joung L.; Holmes, Mark A.; Hong, Xingfang; Mills, Sander INVENTOR (S):

G. :

Parsons, William H.; Sinclair, Peter J.; Steiner. Mark

G.; Wong, Frederick; Zaller, Dennis M. Merck & Co., Inc., USA PCT Int. Appl., 470 pp. CODEN: PIXXD2 Patent PATENT ASSIGNEE(S):

SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT	NO.			KIN		DATE					ION				ATE	
WO 200					-	2001	0104									
W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BĢ,	BR,	BY,	BZ,	CA,	CH,	CN,
	CR,	Cυ,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KR,	ΚŻ,	LC,	LK,	LR,	LS,	LT,	LU,
	LV,	MA,	MD,	MG,	ΜK,	MN,	MW,	ΜX,	ΜŻ,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,
	SE,	SG,	SI,	sĸ,	SL,	TJ,	TM,	TR,	TT,	TZ,	UΑ,	UG,	US,	UΖ,	٧N,	YU,
	ZA,	ZW,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM					
RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,
	DΕ,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ΒJ,
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CA 238																
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EP 120	5265			B1		2003	1112									
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						RO,										
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															0000	

OTHER SOURCE(S): MARPAT 134:86271

L69 ANSWER 19 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) diazabicyclo[3.3.0]oct-3-yl)benzimidazol-1-yl)pyrimidine
R1: RCT (Reactant); SPN (Synchetic preparation); TRU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)
(prepn. of pyrimidine derivs. as Src-family protein tyrosine kinase inhibitor compds.)
RN 317826-43-0 HCAPLUS
CN 2-Pyrimidinamine,
4-[5-(tetrahydro-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)-1H-benzimidazol-1-yl]-N-[(1S)-1-{3-(trifluoromethyl)phenyl]ethyl]- (9CI)
(CA

INDEX NAME)

Absolute stereochemistry.

317826-08-7P, 2-{(s)-1-Phenylethylamino]-4-{5-(1,3-diazabicyclo(3.3.0)oct-3-yl)benzimidazol-1-yl)pyrimidine
317926-09-8P, 2-{(s)-1-Phenylethylamino]-4-{6-(1,3-diazabicyclo(3.3.0)oct-3-yl)benzimidazol-1-yl)pyrimidine
317926-42-9P, 2-{(s)-1-(3-Nitrophenyl)ethylamino]-4-{5-(1,3-diazabicyclo(3.3.0)oct-3-yl)benzimidazol-1-yl)pyrimidine
RL: SPN (Synthetic preparation); TRU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of pyrimidine derivs. as Src-family protein tyrosine

inhibitor compds.) 317826-08-7 HCAPLUS 2-Pyrimidhamine, N-{(IS}-1-phenylethyl]-4-[5-(tetrahydro-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)-1H-benzimidazol-1-yl)- (9CI) (CA INDEX NAME)

(Continued) L69 ANSWER 19 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

317826-09-8 HCAPLUS
2-Pyrimidinamine, N-[(1S)-1-phenylethyl)-4-[6-(tetrahydro-1H-pyrrolo[1,2-cjimidazol-2(3H)-yl)-1H-benzimidazol-1-yl]- (9CI) (CA INDEX NAME)

317826-42-9 HCAPLUS 2-Pyrimidinamine, N-[(1S)-1-(3-nitrophenyl)ethyl]-4-[5-(tetrahydro-1H-pyrrolo(1,2-c]imidazol-2(3H)-yl)-1H-benzimidazol-1-yl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 19 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT: FORMAT

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

L69 ANSWER 20 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 2001:12267 HCAPLUS DOCUMENT NUMBER: 134:71602 Preparation

134:71602
Preparation and effect of benzimidazolylpyrimidine derivatives as SRC kinase inhibitors
Goulet, Joung L.: Holmes, Mark A.; Hunt, Julianne A.;
Mills, Sander G.: Parsons, William H.; Sinclair, INVENTOR (S):

Peter

PATENT ASSIGNEE(S):

J.; Zaller, Dennis M. Merck & Co., Inc., USA PCT Int. Appl., 173 pp. CODEN: PIXXD2 Patent English SOURCE:

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA	TENT	NO.			KIN	0	DATE		- 2	APPL	ICAT	I ON I	NO.		D	ATE	
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WO	2001	0002	07		A1		2001	0104	1	WO 2	000-	US17	510		2	0000	626
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		CR.	cu,	cz.	DE.	DK,	DM.	DZ,	EE,	ES,	FI.	GB,	GD,	GE,	GH,	GM,	HR,
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	pw.				LS.								ZW.	AT.	BE.	CH.	CY.
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	6329																
	1206																
EP																	
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	2003																
RIORIT	Y APP	LN.	INFO	.:						US 1	999-	1416	30P		P 1	9990	630
									,	WO 2	000-	US17	510	,	W 2	0000	626

OTHER SOURCE(S): MARPAT 134:71602

· STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT ·

AB Title Pyrimidine compds: [1; Rl, R2 independently = H, Br, Cl, I, F, OH, SM, CN, NO2, NH2; RlR2; fused methylenedioxy ring, fused 6-membered aromatic ring; Rl, R5 independently = H, alkyl, aryl; RlR5 = O; R4 = H, alkyl, alkoxyl; Xl, X2, X3, X4 independently = CH, CBr, COH, CSH, CNO2, N; R7 = H, NH2, alkyl, aryl, alkylamino, arylamino; Y = O, N, CH, Z = CO, SO2, bond; m, independently = 0, I, 2, 3, 41, or their pharmaceutically acceptable salts, hydrates, solvates, crystal forms and individual disstereomers, and pharmaceutical compns. including the same, which are inhibitors of tyrosine kinase enzymes, and as such are useful in the prophylaxis and treatment of protein tyrosine kinase-associated disorders, such as immune diseases, hyperproliferative disorders and other diseases

L69 ANSWER 20 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) in which inappropriate protein kinase action is believed to play a role, such as cancer, angiogenesis, atherosclerosis, graft rejection, rhewmatoid arthritis and psoriasis. Thus, the title compd. II was prepd. and tested.

IT 315717-68-IP 315717-69-2P
RL: RAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and effect of benzimidazolylpyrimidine derivs. as SRC kinase

se inhibitors)
315717-68-1 HCAPLUS
1-Piperidinecarboxamide, N-phenyl-3-[1-[[4-[5-(tetrahydro-1H-pyrrolo[1,2-c]midazol-2[3H)-yl]-1H-benzimidazol-1-yl]-2-pyrimidinyl]amino]ethyl](9CI) (CA INDEX NAME)

315717-69-2 HCAPLUS
1-Piperidinecerboxamide, N-1-naphthalenyl-3-[1-[[4-[5-(tetrahydro-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)-1H-benzimidazol-1-yl]-2pyrimidinyl]amino]ethyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L69

L69 ANSWER 21 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 2000:139147 HCAPLUS DOCUMENT NUMBER: 132:175859 DOCUMENT NUMBER: TITLE: Drugs containing pyrrolo[1,2-a]pyrazine derivatives ligands for SHTIA receptor and imaging of the organs using the derivatives
Sannar, Mark A.
Pfizer Products Inc., Japan
Jpn. Kokai Tokkyo Koho, 22 pp.
CODEN: JKXXAF
Patent INVENTOR (S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE JP 2000063276 JP 3356726 US 6284757 CA 2280447 CA 2280447 MX 9907598 BR 9906169 A2 B2 B1 AA C A 20000229 JP 1999-230267 19990817 20021216 20010904 US 1999-372438 CA 1999-2280447 19990811 19990813 20000217 20050329 MX 1999-7598 BR 1999-6169 US 1998-96875P 19990817 20000815 PRIORITY APPLN. INFO.: P 19980817

MARPAT 132:175858

OTHER SOURCE(S):

The derivs. I [Rl = Ph, naphthyl, benzoxazolonyl, indolyl, indolonyl, benzimidazolinyl, quinolyl, furyl, benzofuryl, thienyl, benzothienyl, oxazolinyl, benzoxazolyl; R2 = H, Cl-6 alkyl; R3 = Ph, pyridinyl, pyrimidinyl, pyrazinyl, pyridazinyl; R4, R5 = H, Cl-6 alkyl; R1-R3 mes substituted with l-4 f, C, Br, iodo, cyano, NOZ, thiocyano, SR4, SOR4, SO2R4, NHSOR4, Cl-6 alkoxy, NR4R5, NR4COR5, CONR4R5, Ph, COR4, COZR4,

(halo)alkyl, C3-6 cycloalkyl, OCF3; X = 0, S, SO, SO2, NR4, CO, CH(OH), CHR4, CCO, CO2, NR4CO, CONR4; m = 0, l; n = 0, l; l or their pharmaceutically acceptable salts enhance or inhibit serotomergic neurotransmission, and are useful for treatment of diseases, e.g. headache, anxiety, depression, post-traumatic stress disorders, neurodegenerative disorders, prostatic cancer, drug addictions, etc.

claimed are imaging of organs using I labeled with radioisotopes or by combination of I with radiomimetic agents, and compns. for the imaging

are also ligands of dopamine D4 receptor and useful for treatment of

ANSWER 21 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) diseases through enhancing or suppressing dopaminergic neurotransmission. 193068-03-0 arguage=v3-v RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of pyrrolo[1,2-a]pyrazine derivs. as ligands for 5HTIA receptor. receptor)
193068-03-0 HCAPLUS
H-Pyrrolo[1,2-e]imidazole-5,7-dicarboxylic acid, hexahydro-2(phenylmethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

4

L69 ANSWER 22 OF 63 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: octabud-HCAPLUS COPYRIGHT 2006 ACS on STN 1998:98051 HCAPLUS 128:154101

octahydropyrrolo[1,2-

creparation of 2,7-disubstituted

alpyrazine derivatives as dopamine D4 receptor ligands.
Sanner, Mark A.
Pfizer Inc., USA
U.S., 26 pp.
CODEN: USXXAM
Patent
English
1 INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE US 5714487 PRIORITY APPLN. INFO.: US 1996-774290 US 1996-774290 19961223 19961223 19980203

OTHER SOURCE(S): MARPAT 128:154101

Title compds. [I; R1 = Ph, naphthyl, benzoxazolonyl, indolyl, indolonyl, benzimidazolyl, quinolyl, furyl, benzofuryl, thienyl, benzoxaclyl, oenzoxaclyl, benzoxaclyl; R2, R4 = H, alkyl: R3 = Ph, pyridyl, pyrimidinyl, pyrazinyl, pyridzinyl; m, n = 0-2; X = O, S, SO, SO2, NR4, CO, CHOH,

CHR4, CONR4, etc.], were prepared Thus, (7RS,8aSR)-7-(4-

fluorophenoxy)methyl-2-phenylmethyl-1,2,3,4,6,7,8,8a-octahydropyrrolo[1,2-a]pyrazine (preparation given) was refluxed with ammonium formate and Pd/C in

in MeOH and the residue was refluxed with 2-chloro-6-fluoropyrimidine and Na2CO3 in H2O to give (7RS,8aSR)-7-(4-fluorophenoxy)methyl-2-(5-fluoropyrimidin-2-yl)-1,2,3,4,6,7,8,8a-octahydropyrrolo[1,2-a]pyrazine.

I showed binding affinities for displacement of [3H]-spiperone of <2 μM.

17 193058-03-0
R1: RCT (Reactant): RACT (Reactant or reagent)
(preparation of 2,7-disubstituted octahydropyrrolo[1,2-a]pyrazine
derivs. as
dopamine D4 receptor ligands)
RN 193068-03-0 HCAPEUS
CN H-Pyrrolo[1,2-c]imidazole-5,7-dicarboxylic acid, hexahydro-2(phenylmethyl)-, dimethyl ester (9CI) (CA INDEX NAME)

L69 ANSWER 22 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT: THIS

FORMAT

THERE ARE 11 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L69 ANSWER 23 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1997:753972 HCAPLUS DOCUMENT NUMBER: 128:123431

1991:7339/2 nurrows 128:122431 Inhibition of frog skeletal muscle sodium channels by newly synthesized chiral derivatives of mexiletine TITLE:

CORPORATE SOURCE:

tocainide
De Luca, Annamaria; Natuzzi, Fedele; Falcone, Giulia;
Duranti, Andrea; Lentini, Giovanni; Franchini, Carlo;
Tortorella, Vincenzo; Conte Camerino, D.
Facolta di Farmacia, Dipartimento Farmacobiologico,
Unita di Farmacologia, Via Orabona 4, Bari, I-70125,
Italy
Naunyn-Schmiedeberg's Archives of Pharmacology AUTHOR (S) :

(1997),

356(6), 777-787

CODEN: NSAPCC; ISSN: 0028-1298

PUBLISHER: Springer-Verlag

DOCUMENT TYPE: Journal

LANGUAGE: English

AB To search for potent use-dependent blockers of skeletal muscle sodium

channels as potential antimyotonic agents, the actions of newly

synthesized chiral analogs of mexiletine and tocainide were tested in

vitro on sodium currents of single fibers of frog semitendinosus muscle

by

bу vaseline-gap voltage clamp method. The effect of each drug on the

maximal ' peak Na+ transient (INa max) was evaluated as both tonic and

use-dependent block by using infrequent depolarizing stimulation and trains of pulses at

at

2-10 Hz frequency, resp. The mexiletine analog

3-(2,6-dimethylphenoxy)-2methylpropanamine (Me2), having an increased distance between the Ph and
the amino groups, was less potent than mexiletine in producing a tonic
block but produced a remarkable use-dependent block. In fact, the
half-maximal concentration (ICSO) for tonic block of S(-)-Me2 was 108

half-maximal concentration (ICSO) for tonic block of S(-)-Me2 was 108 µM vs.

54.5 µM of R(-)-mexiletine, but the ICSO was 6.2 times lowered by the
10 Hz stimulation with respect to the 2.4-fold decrease observed with
mexiletine. The R(-)-mexiletine and the S(-)-Me2 were about twofold more
potent than the corresponding enantiomers in producing a tonic block, but
the stereoselectivity attenuated during use-dependent blockade. The more
lipophilic 2-(4-chloro-2-methylphenoxy)-1-phenylethylamine (Me1),
presently available as raceme, produced a potent and irreversible tonic
block of the sodium currents with an ICSO of 29 µM, but had a less
pronounced use-dependent inhibition, with a 1.9-fold decrease of the ICSO
at 10 Hz. The R(-) isomer of 2',6'-valinoxylidide (Tol), a tocainide
derivative with an increased hindrance on the chiral carbon atom, was
twofold

derivative with an increased ninorance on the chiral carbon atom, was twofold (IC50 = 209 µM) and tenfold (IC50 = 27.4 µM) more potent than R(-)-tocainide in tonic and use-dependent block, resp. Tocainide was almost devoid of stereoselectivity, whereas the eudismic ratio of Tol [IC50 S(+)-Tol/IC50 R(-)-Tol] was 1.7. As for mexiletine and Me2, the stereoselectivity of Tol was the weaker the higher the frequency of stimulation. The cyclic pyrroloimidazolonic tocainide analog To2

L69 ANSWER 23 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

● HBr

REFERENCE COUNT: THERE ARE 36 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

ANSWER 23 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) a small tonic block at 500  $\mu M_{\odot}$  and 1 min stimulation at 10 Hz was needed to show up a 50 block of INa max. All the compds. produced a left-shift of the steady-state inactivation curve correlated pos. wi

extent of use-dependent inhibition, with the exception of the cyclic To2 that acted as an open-channel blocker. The highly use-dependent blockers Me2 and Tol might be promising drugs to solve high frequency discharges

of

action potentials typical of myotonic muscles. Concomitantly the high
potency of Mel and the open-channel block exerted by To2 can represent
important features to get selective blockers for skeletal muscle sodium
channels.

IT 201986-87-0 201986-88-1
RL: BaC (Biological activity or effector, except adverse); BSU
(Biological
study, unclassified); BIOL (Biological study)
(inhibition of frog skeletal muscle sodium channels by newly
synthesized chiral derivs. of mexiletine and tocainide)

RN 201986-87-0 HCAPLUS
CN 1H-Pyrrolo(1,2-c]mida2o1-1-one, 2-(4-chlorophenyl)hexahydro-,
monohydrobromide, (R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

• HBr

201986-88-1 HCAPLUS 1H-Pyrrolo(1,2-c)imidazol-1-one, 2-(4-chlorophenyl)hexahydro-, monohydrobromide, (5)- (9CI) (CA INDEX NAME)

L69 ANSWER 24 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1997:528590 HCAPLUS DOCUMENT NUMBER: 127:130461 TITLE: 5ynthesis and Structure-Activity

1397:132830 NACHUS
127:130461
Synthesis and Structure-Activity Relationships of a New Model of Arylpiperazines. 3. 2-[a-(4-Arylpiperazin-1-yl)alkyl]perhydropyrrolo[1,2-c]imidazol s and -perhydroimidazol[,5-a]pyridines: Study of the Influence of the Terminal Amide Fragment on 5-HTIA Affinity/Selectivity
Lopez-Rodriguez, Maria L.: Morcillo, M. Jose;
Fernandez, Esther: Porras, Esther: Murcia, Marta;
Sanz, Antonio M.: Orensanz, Luis
Departamento de Quimica Organica I Facultad de Ciencias Quimicas, Universidad Complutense, Madrid, 28040, Spain
Journal of Medicinal Chemistry (1997), 40(16), 2653-2656
CODEN: JMCMAR; ISSN: 0022-2623

AUTHOR (S):

CORPORATE SOURCE:

CODEN: JMCMAR; ISSN: 0022-2623 American Chemical Society

PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

MENT TYPE: Journal
UNGE: English
A series of new arylpiperazine derivs., which are devoid of the terminal
amide fragment present in related 5-HTIA ligands, was prepared and

wated for affinity at 5-HTIA and  $\alpha$ l receptors. All the compds. demonstrated high affinity for the 5-HTIA receptor and moderate affinity for  $\alpha$ l receptor binding sites. Structure-activity relationship (SAR) studies suggest that there is influence of electronic factors on

the no-pharmacophoric part of the dl receptor site. However there is no influence of electronic interactions on the stabilization of the 5-HTIA receptor-ligand complex. 12293-09-19 RL: BPR (Biological process); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation);

(Process)

SOURCE:

(Process)
(preparation and affinity at 01- and 5-HTIA-receptors of
arylpiperazines)
19299-08-1 HCAPLUS
1H-Pytrolo[1,2-c]imidazole, hexahydro-2-[3-[4-(2-methoxyphenyl)-1-piperazinyl]propyl]-, tetrahydrochloride (9CI) (CA INDEX NAME)

●4 HC1

192992-82-89 192992-83-99 192992-86-29 192992-87-39 192992-88-49 RI: BPR (Biological process); BSU (Biological study, unclassified); SPN

ANSWER 24 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
(Synthetic preparation); THU (Therapeutic use); BIOL (Biological study);
PREP (Preparation); PROC (Process); USES (Uses)
(prepn. and affinity at al- and 5-HT1A-receptors of
arylpiperazines)
192992-82-8 HCAPLUS
HI-Pyrrolo[1,2-c|imidazole, hexahydro-2-(3-[4-(2-methoxyphenyl)-1-piperazinyl)propyl]- (9CI) (CA INDEX NAME)

RN 192992-83-9 HCAPLUS
CN 1H-Pyrrolo(1,2-c)imidazole,
hexahydro-2-[3-{4-[3-(trifluoromethyl)phenyl}l-piperazinyl]propyl]- (9CI) (CA INDEX NAME)

192992-86-2 HCAPLUS
1H-Pyrrolo[1,2-c]imidazole, hexahydro-2-[4-[4-{2-methoxyphenyl}-1-piperazinyl]butyl]- (9CI) (CA INDEX NAME)

(CH<sub>2</sub>)<sub>3</sub>-

192992-87-3 HCAPLUS
1H-Pyrrolo[1,2-c]imidazole, 2-[4-[4-(3-chlorophenyl)-1-piperazinyl]butyl]hexahydro- (9CI) (CA INDEX NAME)

L69 ANSWER 25 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1997:506300 HCAPLUS DOCUMENT NUMBER: 127:135811

127:135811
Preparation of 2,7-substituted octahydropyrrolo[1,2-a)pyrazine derivatives as ligands for dopamine receptor subtypes
Sanner, Mark A.
Pfizer Inc., USA; Sanner, Mark A.
PCT Int. Appl., 73 pp.
CODEN: PIXXD2 TITLE:

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: Patent English LANGUAGE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PAT	TENT	NO.			KIN	)	DATE			APP	LICA	TION	NO.		D	ATE	
													192		1	9961	106
													, KR,				
													, UZ,				
	RW:	AT.	BE.	CH.	DE.	DK.	ES,	FI.	FR.	GB	. GR	. IE	, IT,	LU,	MC,	NL,	PT,
		er.	DF	D.T	CF	CC	CT	CM	Ch	CN	MT.	MD	NE	QM.	тb	TG	
CA	2240	594			AA		1997	0703		CA	1996	-224	0594 80 226		1	9961	106
CA	2240	594			C		2001	0724									
AU	9673	280			A1		1997	0717		ΑU	1996	-732	80		1	9961	106
AU	7045	78			B2		1999	0429									
EP	8748	49			A1		1998	1104		EΡ	1996	-935	226		1	9961	106
EP	8748	49			В1		2001	0919									
	R:	AT.	BE.	CH.	DE.	DK.	ES.	FR.	GB.	GR	. IT	, LI	, LU,	NL.	SE,	PT,	IE,
CN	1205	704			А		1999	0120		CN	1996	-199	250 46 446 742		1	9961	106
CN	1061	350			В		2001	0131									
BR	9612	246			А		1999	0713		BR	1996	-122	46		1	9961	106
JP	1150	8920			T2		1999	0803		JΡ	1997	-523	446		1	9961	106
JΡ	3204	456			B2		2001	0904									
RU	2162	470			C2		2001	0127									
AT	2058	46			E		2001	1015		AT	1996	-935	226		1	9961	106
ES	2161	377			<b>T</b> 3		2001	1201		ES	1996	-935	226		1	9961	106
PT	8748	49			т		2002	0130		PT	1996	-935	226		1	9961	106
TW	2161 8748 4790	58			В		2002	0311		TW	1996	-851	13669		1	9961	108
										ZA	1996	-107	81 3		1	9961	220
NO	9802 3099	843			А		1998	0619		NO	1998	-284	3		1	9980	619
NO	3099	36			В1		2001	0423									
GR	3037	060			Т3		2002	0131		GR	2001	-401	932		2	0011	
	APP									US	1995	-898	8 P		P 1	9951	221
													192				

OTHER SOURCE(S): MARPAT 127:135811

ANSWER 24 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN 192992-88-4 HCAPLUS 1H-Pyrrolo[1,2-c]imidazole, hydro-2-[4-[4-[3-(trifluoromethyl]phenyl]-1-piperazinyl]butyl]- (9CI) (CA INDEX NAME) (Continued)

REFERENCE COUNT:

THERE ARE 34 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L69 ANSWER 25 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) quinolyl, furyl, benzofuryl, thienyl, benzothienyl, oxazolyl, benzoxazolyl; R2 = H, (C1-C6)alkyl; R3 = Ph, pyridinyl, pyrimidinyl, pyrazinyl, pyridazinyl; X = O, S, OS, Ox, NAR, CO, CH(OH), CRR4, etc.; m = 0, 1, 2; n = 0, 1, 2] were prepd as ligands for dopamine receptor subtypes, esp. the dopamine D4 receptor. E.g., (7Rs,8aSR)-7-(4-fluorophenoxy)-2-phenylmentyl-1,2,3,4,6,7,8,8a-octahydropyrrolo1,2-alpyrazine and aq. ammonium formate in MeOH was treated with an aq. slurry

a)pyrazine and aq. ammonium rotation and an ammonium rotation of 10% Pd/C and the product then reacted with 2-chloro-5-fluoropyrimidine to give (7RS, 88R)-7-(4-fluorophenoxy)-2-(5-fluoropyrimidin-2-yl)-1,2,3,4,6,7,8,8a-octahydropyrrolo(1,2-a)pyrazine. The title compds. had binding affinities for the displacement of [3H]-spiperone < 2

D4 binding affinities for the displacement of [3H]-spiperone < 2 micromolar.

IT 193068-03-0
RL: RCT (Reactant): RACT (Reactant or reagent) (preparation of 2,7-substituted octahydropyrrolo[1,2-a]pyrazine derivs. as ligands for D4 dopamine receptors)
RN 193068-03-0 HCAPLUS
CN 1H-Pyrrolo[1,2-c]midazole-5,7-dicarboxylic acid, hexahydro-2-(phenylmethyl)-, dimethyl ester (9CI) (CA INDEX NAME)



L69 ANSWER 26 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1996:701300 HCAPLUS DOCUMENT NUMBER: 126:89181

DOCUMENT NUMBER:

126:89181
Synthesis and antibacterial activity of 1-B-methylcarbapenems having a 1,3-diazabicyclo{3.3.0]octan-4-one moiety Nam, Ki Hong: Oh, Chang Hyunc Cho, Jin Koo: Kim, Hyo Jung: Lee, Ki Soo: Cho, Jung Hyuck Division Applied Science, Korea Institute Science Technology, Seoul, 130-650, S. Korea Archiv der Pharmazie (Welnheim, Germany) (1996), 329(10), 443-446
CODEN: ARPMAS: ISSN: 0365-6233
VCH
JOURNAL FORMARY CONTRACTOR OF THE PROPERTY OF T AUTHOR (S):

CORPORATE SOURCE:

SOURCE:

PUBLISHER MENT TYPE:

DOCUMENT LANGUAGE:

The synthesis of the methylcarbapenems I (R, Rl = H; R = Me; Rl = H, Me, £t, Pr, Ph or R = cyclopropyl; Rl = H, Me) from protected 2-(diphenylphosphocyloxy) carbapenem and the appropriate mercaptoethyldiazabicyclocetanone is described. Their in-vitro antibacterial activities against both Gram-pos. and Gram-neg. bacteria

are

are reported. The effect of the substituent on the bicyclic ring was investigated in agreement with findings from our previous studies.

IT 185736-63-49

RE: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study, PREP (Preparation)

(preparation and antibacterial activity of diazabicyclooctanone-substituted activity of diazabicyclooctanone-substituted carbapenems)

RN 185736-63-4 HCAPLUS

RN 185736-63-4 HCAPLUS

CN 1-Azabicyclo(3.2.0)hept-2-ene-2-carboxylic acid, 6-(1-hydroxyethyl)-4-methyl-7-oxo-3-[[2-(tetrahydro-1-oxo-1H-pyrrolo[1,2-c]nmidazol-2(3H)-yllethyl]thio]-, [4R-[3(S\*),4a,5B,6β(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 26 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

185736-80-5
RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation and antibacterial activity of
zabicyclooctanone-substituted
carbapenems)
185736-80-5 HCAPLUS
1-Azabicyclo[3.2.0]hept-2-ene-2-carboxylic acid, 6-(1-hydroxyethyl)-4methyl-7-oxo-3-[{2-(tetrahydro-1-oxo-1H-pyrrolo[1,2-c)imidazol-2(3H)yl)ethyl|thio|-, (4-nitrophenyl|methyl ester, [4R[3(S\*),4α,5β,6β(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 27 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1996:674366 HCAPLUS DOCUMENT NUMBER: 125:322383 TITLE: Preparation

Preparation of novel carbapenem derivatives as antibacterials

INVENTOR (S):

antibacterials
Aihara, Kazuhiro; Kano, Yuko; Shiokawa, Sohjiro;
Sasaki, Toshiro; Setsu, Fumihito; Toyooka, Yumiko;
Ishil, Miyuki; Atsumi, Kunio; Iwamatsu, Katsuyoshi;
Tamura, Atsushi
Melji Seika Kabushiki Kaisha, Japan
PCT Int. Appl., 107 pp.
CODEN: PIXXD2
Patent

PATENT ASSIGNEE (S): SOURCE:

DOCUMENT TYPE: Patent Japanese

	PA:	PENT	NO.					DATE			API	LI	CAT	ION	NO.			ATE	
	WO	9628	455			A1		1996	0919		wo	19	96-	JP57	3			9960	
								KR.	PL.	SI,	US	;	-						
		RW:	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR,	GE	3,	GR,	IE,	IT,	LU,	MC,	NL,	PT,
SE																			
	CA	2189	995			AA		1996	0919		CA	19	96-	2189	995		1	9960	308
	CA	2189	995			С		2001	0123										
	ΕP	7603	70			A1		1997	0305		EΡ	19	96-	9050	36		1	9960	308
	EP	7603	70			B1		2002	0807										
		R:	BE,	DE,	ES,	FR,	GB,	IT,											
	CN	1148	390						0423		CN	19	96-	1901	77		1	9960	308
		1057				В			1004										
		2179				Т3			0201						36			9960	
		4253				В			0311						2872			9960	
		5990				А		1999	1123						32			9970	
PRIC	RIT	Y APP	LN.	INFO	.:						JP	19	95-	5161	6	i	A 1	9950	310
															-				

OTHER SOURCE(S): MARPAT 125:328383

Title compds. I [R1 = H, alkyl, R2-R5 = H, halo, OH, nitro, cyano, COOH, formyl, alkyl, cycloalkyl, C2-4 alkenyl, C2-4 alkynyl, alkoxy, etc.] are prepared The compds. have a broad and potent antibacterial activity on Gram-pos. bacteria and Gram-neg, bacteria including Pseudomonas uginosa and show a potent antibacterial effect on various  $\beta$ -lactamase-producing bacteria and MRSA and an extremely high DHP-1 stability. Thus,

L69 ANSWER 27 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

allyl
{Is,5R,6S}-6-[(IR)-1-(allyloxycarbonyloxy)ethyl]-2-(hydroxymethyl)-1methyl-1-carbapen-2-em-2-carboxylate was reacted with di-Ph
phosphorochloridate in CH2Cl2 contg. 4-(dimethylamino)pyridine to give

corresponding phosphate, which was reacted with 3(hydroxymethyl)imidazo[5,1-b]thiazole in DMF contg. NaI, and the product
treated with Ph3P, 2-ethylhexanoic acid, potassium 2-ethylhexanoate, and
tetrakis(triphenylphosphine)palladium in (H2C12 at room temp. for 2 h to
give the title compd. I [R1 = Me, R2 = CH2OH, R3-R5 = H]. This had an

give the title compd. 1 [R] = Me, RZ = CHZOH, R3-R5 = H]. This had an MIC comparable to that of imipenem/cilastatin against Staphylococcus aureus. Pharmaceutical compns. contg. I are described.

IT 183067-38-1P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study; except activity or effector, except adverse); BSU (Biological study); PREP (Preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of novel carbapenem derivs. as antibacterials)

RN 183067-38-1 HcAPLUS
CN 5H-Cyclopent(d|imidazo[5,1-b|thiazolium, 2-[[2-carboxy-6-(1-hydroxyethy])-4-methyl-7-oxo-1-azabicyclo[3,2.0]hept-2-en-3-yl]methyl]-6,7-dihydro-, inner salt, [4S-[4a,5],6](S+)]]- (SCI (CA INDEX NAME)

L69 ANSWER 28 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1995:1000583 HCAPLUS
DOCUMENT NUMBER: 124:175947
TITLE: An efficient construction of 4-oxo-1,3-diszabicyclo[3,3,0] octanes via thiohydantoins
AUTHOR(S): Kim, In Jong; Yoo, Kyung Ho; Shin, Kye Jung; Kim,

AUTHOR (S): Dong

Jin; Park, Sang Woo Div. Applied Science, Korea Inst. Science Technology, Seoul, 131-650, S. Korea Synthetic Communications (1995), 25(24), 4001-10 CODEN: SYNCAV; ISSN: 0039-7911 Dekker Journal English CASREACT 124:175947 CORPORATE SOURCE:

SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

OTHER SOURCE(S):

New stereoisomeric N-bridged heterocycles, 4-oxo-1,3-dlazabicyclo $\{3.3.0\}$  octanes  $\{I:R=Me,Et,Ph,4-MecSH4,X=H,H\}$  were synthesized from trans-4-hydroxy-L-proline  $\{II\}$ . Thiohydantoins I  $\{X=Me,H\}$ 

as the key intermediates were prepared by nucleophilic addition of II to isothiocyanates, and subsequent cyclization. These thiohydantoins I (X = S) were readily desulfurized to provide I (X = H,H). 173549-72-99 173549-73-0P 173549-74-1P 173549-75-2P 173658-16-7P 173658-17-8P 173658-18-9P 173658-19-0P RESPONDED TO THE PROPERTY OF TH

thiohydantoins)
173549-72-9 HCAPLUS
1H-Pytrolo[1,2-c]imidazol-1-one, hexahydro-6-hydroxy-2-methyl-, (6R-cis)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 28 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

173658-17-8 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, 2-ethylhexahydro-6-hydroxy-, (6R-trans)-(9C1) (CA INDEX NAME)

Absolute stereochemistry.

173658-18-9 HCAPLUS 1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-6-hydroxy-2-phenyl-, (6R-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

173658-19-0 HCAPLUS |H-Pyrrolo[1,2-c|limidazol-1-one, hexahydro-6-hydroxy-2-(4-methylphenyl)-, (6R-trans)- (9C1) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 28 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) 173549-73-0 HCAPLUS H-Pyrrolo(1,2-c)imidazol-1-one, 2-ethylhexahydro-6-hydroxy-, (6R-cis)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

173549-74-1 HCAPLUS
HH-Pyrrolo(1,2-c)imidazol-1-one, hexahydro-6-hydroxy-2-phenyl-, (6R,7aS)-(SCI) (CA INDEX NAME)

Absolute stereochemistry.

173549-75-2 HCAPLUS 1H-Pyrrolo(1,2-c|imidazol-1-one, hexahydro-6-hydroxy-2-(4-methylphenyl)-, (6R-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

173658-16-7 HCAPLUS HH-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-6-hydroxy-2-methyl-, (GR-trans)- (SCI) (CA INDEX NAME)

Absolute stereochemistry



L69 ANSWER 29 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1995:713780 HCAPLUS DOCUMENT NUMBER: 123:111745
TITLE: Preparation of antibacterial cell inventor(S): Acsumi, Kunio; Umemura, Eijiro; Cabibilia Number (S): Cabibilia Numb PARLUS
123:111745
Preparation of antibacterial cephem derivatives
Atsumi, Kunio; Umemura, Eijiro; Kano, Yuko; Shiokawa,
Sohjiro; Kudo, Toshinaki; Tsushima, Masaki; Iwamatsu,
Katsuyoshi; Tamura, Atsushi; Shibahara, Seiji
Meiji Seika K. K., Japan
PCT Int. Appl., 326 pp.
CODEN: PIXXD2
Patent
Japanese
1

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	TENT 1						DATE		AP	PL	ICAT	ION	NO.			DATE	
						-											
WO	9507	912			A1		1995	0323	WO	1	994-	JP1	529			19940	916
	W:	CA,	CN,	JP,	KR,	US											
	RW:	AT,	BE,	CH,	DE,	DK	, ES,	FR,	GB, G	R,	IE,	IT	, LU,	MC,	NL	, PT,	SE
EP	6693	36			A1		1995	0830	EP	1:	994-	927	055			19940	916
EP	6693	36			В1		2000	0517									
	R:	AT,	BE,	CH,	DE,	ES	, FR,	GB,	IE, I	Т,	LI,	NL					
CN	1114	507			A		1996	0103	CN	1	994-	190	696			19940	916
CN	1046	286			В		1999	1110									
TW	3853	12					2000	0321	TW	1	994-	831	08591			19940	916
	19292						2000	0615	AT	1	994-	927	055			19940	916
	2146						2000	0816	ES	1	994-	927	055			19940	916
CA	2149	514			С		2000	1031	CA	. 1	994-	214	9514			19940	916
JP	3152	934			B2		2001	0403	JP	1	995-	509	092			19940	916
US	5663	162			A		1997	0902	US	1	995-	436	280			19950	725
PRIORIT	Y APP	LN.	INFO	.:					JP	1	993-	230	573	,	A.	19930	916
									JP	1	994-	211	908	,	A.	19940	812
									WO	1	994-	JP1	529	١	a	19940	916

OTHER SOURCE(S): MARPAT 123:111745

Title compds. I [R1 represents H, alkyl, alkenyl, etc.; and R2, R3, R4

L69 ANSWER 29 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) 2-methoxyiminolacetamido-3-(chloromethyl)-3-cephem-4-carboxylic acid p-methoxybenzyl ester was reacted with imidazo[5,1-b]thiazole (prepn given) in acetone conto. NaI at room temp. overnight to give, after hydrolysis, (68,78)-7-[(2)-2-(2-aminothiazol-4-yl)-2-

methoxyiminoacetamido]-3-(imidazo[5,1-b]thiazolium-6-yimethyl)-3-cephem-4-carboxylate inner salt. (6R,7R)-7-(Z)-2-(2-aminothiazol-4-yl)-2-([5]-1-carboxyethoxyiminoacetamido]-3-(imidazo[5,1-b]thiazolium-6-ylmethyl)-3-cephem-4-carboxylate (also prepd.) had an MIC of 6.25 µg/mL against Staphylococcus aureus. Pharmaceutical compns. contg. I are described.

IT 165665-19-0P 165665-20-3P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological

logical study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of antibacterial cephem derivs.) 163655-19-0 HCAPLUS (Preparation); Polythiazolium, 2-[(7-[(2-amino-4-thiazoly!)(methoxyimino)acety!)amino|-2-carboxy-8-oxo-5-thia-1-azabicyclo[(4.2.0)oct-2-en-3-yl]methyl]-6,7-dihydro-, inner salt, [6R-[6a,78(2)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

RN 165665-20-3 HCAPLUS
CN 5H-Cyclopent[d]imidazo[5,1-b]thiazolium,
2-[{T-[[2-amino-4-thiazoly]][[1-carboxyethoxy]imino]acety]]amino]-2-carboxye-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-an-3-yl]methyl]-6,7-dihydro-, inner salt,
[6R-[6\(\alpha\),7\(\beta\)[2]-(S^1)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

L69 ANSWER 29 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

Absolute stereochemistry.

Absolute stereochemistry.

L69 ANSWER 30 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

108309-32-6 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4thia-1-azabicyclo[3,2,0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner
salt, [5R-[3(R\*),5c,6c(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

RN 118776-90-2 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
2-{2-amino-2-oxoethyl)-6-[[5R,6S)-2-carboxy6-[(1R)-1-hydroxyethyl]-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3yl]thio]-6,7-dihydro-, inner salt (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 30 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

157683-42-6 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-ethyl-6,7-dihydro-, innersalt, [5R-[3(R\*),5a,6u(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

157683-43-7 HCAPLUS  $5N-Pyrrolo[1,2-c]imidazolium, 6-{[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl}thio]-2-ethyl-6,7-dihydro-, inner salt, <math>[5R-[3(s^*),5\alpha,6\alpha(R^*)]]-$  (9CI) (CA INDEX NAME)

Absolute stereochemistry.

157683-50-6 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2,0]hept-2-en-3-y][thio]-6,7-dihydro-2-[2-(methylamino)-2-oxoethyl]-, inner salt, [5R-[3(R\*),5a,6a(R\*)]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 30 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

157683-54-0 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-y1]thio]-6,7-dihydro-2-(phenylmethy1)-, inner salt, [5R-[5 $\alpha$ ,6 $\alpha$ (R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 30 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

157683-51-7 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[2-(methylamio-2-oxo-ethyl]-, inner salt, [5R-[3(8\*),5o,6a(R\*)]]-[9CI] (CA INDEX NAME)

Absolute stereochemistry.

157683-52-8 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 6-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-2-[2-(dimethylamino)-2-oxoethyl]-6,7-dihydro-, inner salt, [5R-[3[R\*], 5a,6a[R\*])]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

157683-53-9 HCAPLUS  $5H-Pyrrolo[1,2-c]imidazolium, \ 6-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-[2-(dimethylamino)-2-oxoethyl]-6,7-dihydro-, inner salt, [5R-[3(S*),5\alpha,6\alpha(R*)]]-(9CI) (CA INDEX NAME)$ 

Absolute stereochemistry.

L69 ANSWER 31 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1994:483169 HCAPLUS
DOCUMENT NUMBER: 121:83169
TITLE: Synthesis of new quinolinone antibacterial agents
with

Characteristics of the Control of the Control of Characteristics of Ch CORPORATE SOURCE:

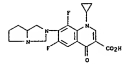
SOURCE:

DOCUMENT TYPE: LANGUAGE: GI

The title compds., e.g., I and II, were prepared by reaction of 7-haloquinolinones with heterocycles. The antibacterial activity of the products was lower than that of ciprofloxacin. 156171-70-99

IT 156171-70-99
RL: SPM (Synthetic preparation); PREP (Preparation)
(preparation of)
RN 156171-70-9 HCAPLUS
CN 3-Quinolinecarboxylic acid,
1-cyclopropyl-6,8-difluoro-1,4-dihydro-4-oxo-7(tetrahydro-1H-pyrrolo(1,2-c)imidazo1-2(3H)-yl)- (9CI) (CA INDEX NAME)

L69 ANSWER 31 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)





L69 ANSMER 32 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1994:116630 HCAPLUS
DOCUMENT NUMBER: 120:116630
TITLE: 4DTHOR(S): H- and 13C-NVR studies of aminoglycoside antibiotics
AUTHOR(S): Moloney, Gerard P.; Craik, David J.; Iskander, Magdy
N.

CORPORATE SOURCE: Victorian Coll. Pharm., Monash Univ., Parkville,

3052,

Australia
SOURCE: Magnetic Resonance in Chemistry (1993), 31(12), 1077-84

CODEN: MRCHEG; ISSN: 0749-1581

JOURNAL JOUR

reported and compared with previous results for the related antibiotic lincomycin. The stability of the 2 cyclized derivs. in aqueous solns.

examined Both cyclizations involved formation of a 4-imidazolidinone

examined Both cyclizations involved formation of a normalization of a normalization.

The ring system based on cyclization with formaldehyde was stable in aqueous solution, whereas that based on benzaldehyde was not.

IT 35119-67-6

RL: PRP (Properties)

(conformation of, proton and carbon-13 NMR study of, stability in relation to)

RN 35119-67-6 HCAPLUS

CN L-chreo-a-D-galacto-Octopyranoside, methyl 7-chloro-6,7,8-trideoxy-6
(tetrahydro-1-oxo-6-propyl-1H-pyrrolo[1,2-c]imidazol-2(3H)-y1)-1-thio-,

(6R-cia)- (9CI) (CA INDEX NAME)

L69 ANSWER 33 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1992:407730 HCAPLUS
DOCUMENT NUMBER: 17:7730 HCAPLUS
ITILE: Preparation of carbapenem derivatives
INVENTOR(S): Suzaki, Hiroshi: Nishi, Toshiyuki: Takemura, Makoto: Hayano, Takeshi
PATENT ASSIGNEE(S): Daiichi Seiyaku Co., Ltd., Japan
Jpn. Kokai Tokkyo Koho, 11 pp.
CODEN: JKXXAF
DOCUMENT TYPE: Patent
LANGUAGE: PAHLLY ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PRIC

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 04009380	A2	19920114	JP 1990-109307	19900425
JP 3045518	B2	20000529		
ORITY APPLN. INFO.:			JP 1990-109307	19900425
FR SOURCE (S) .	маррат	117-7730		

OTHER SOURCE(S): MARPAT 117:7730

STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

Carbapenem derivs. [I: R1 = alkyl, (protected) hydroxyalkyl; R2 = H, protecting group, anion; R3 = H, alkyl; R4 = (aubstituted) fused heterocyclyl containing  $\geq$ 2 N atoms and an onium center], useful as antibacterial agents, are prepared MeI was added to a solution of 165

mg eater
II in Me2CO with stirring at 5°, more MeI was added, and the mixture
was stirred at 5° and the distillate residue was dissolved in
phosphate buffer and hydrogenolyzed over 10% Pd-C at 4 atm H to give 23

(1R,5S,6S,8R)-III (IV) and 18 mg isomer. IV showed MIC of <0.1 µg/mL against Eacherichia coli NiHJ, etc.
118776-50-4P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and reaction of, in preparation of antibacterial agent)
118776-50-4 HCAPLUS
5H-Pyrrolo(1,2-c]imidazolium, 2-(2-amino-2-oxoethyl)-6,7-dihydro-7-mercapto-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 118776-49-1 CMF C8 H12 N3 O S

ANSWER 33 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

2

IT 141547-45-7P 141547-46-8P 141547-47-9P
141611-04-3P 141611-05-4P 141611-06-5P
141611-07-9F 141611-08-7P
RI: BAC (Biological activity or effector, except adverse); BSU
(Biological
study, unclassified); SPN (Synthetic preparation); BIOL (Biological
study); PREP (Preparation)
(preparation of, as antibacterial agent)
RN 141547-45-7 HCAPMUS
CN 5H-Pyrrolo[1,2-c]imidazolium, 2-(2-amino-2-oxoethyl)-7-[(2-carboxy-6-(1-hydroxyethyl)-4-methyl-7-oxo-1-azabicyclo[3,2,0]hept-2-en-3-yl]thio]-6,7-dihydro-, inner salt, [4R-[3(5\*), 4α,5β,6β(R\*)]]- (9CI)

Absolute stereochemistry.

RN 141547-47-9 HCAPLUS CN 5H-Pyrrolo[1,2-c]imidazolium, 7-{[2-carboxy-6-(1-hydroxyethyl)-4-methyl-7-

oxo-1-azabicyclo(3.2.0]hept-2-en-3-yl}thio]-6,7-dihydro-2-{2-(methylamino)2-oxoethyl]-, inner salt, {4R-{3(S\*),4α,5β,6β(R\*)}}- (9CI)
(CA INDEX NAME)

Absolute stereochemistry

RN 141611-04-3 HCAPLUS
CN 5H-Pyrcolo[1,2-c]imidazolium,
7-[{2-carboxy-6-(1-hydroxyethyl)-4-methyl-7oxo-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner
salt, [4R-[3(S\*), 4α,5β,6β(R\*)])- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 33 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) Absolute stereochemistry.

IT 141547-44-6
RL: RCT (Reactant); RACT (Reactant or reagent)
{reaction of, with carbapenem derivative, in preparation of antibacterial agent)
RN 141547-44-6 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium, 2-ethyl-6,7-dihydro-7-mercapto-, salt with trifluoromethaneaulfonic acid (1:1) (9CI) (CA INDEX NAME)

CRN 141547-43-5 CMF C8 H13 N2 S

CM 2 CRN 37181-39-8 CMF C F3 03 S

L69 ANSWER 33 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN Absolute stereochemistry. (Continued)

141611-06-5 HCAPLUS 5H-Pytrolo[1, 2-c]imidazolium, 2-(2-amino-2-oxoethyl)-7-[[2-carboxy-6-(1-hydroxyethyl)-4-methyl-7-oxo-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-, inner salt, [4R-[3(R\*),  $\{\alpha, 5\beta, 6\beta(R^*)\}]$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

RN 141611-08-7 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
7-[[2-carboxy-6-(1-hydroxyethy1)-4-methy1-7-

oxo-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[2-(methylamino)-2-oxoethyl]-, inner salt,  $\{4R-\{3(R^*),4\alpha,5\beta,6\beta(R^*)\}\}$ - (9CI) (CA INDEX NAME)

L69 ANSWER 34 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
115:114421 HCAPLUS
115:114421
Heterocyclization of the 2-aminoalkyl(and aryl)benzimidazoles under phase transfer catalysis conditions
AUTHOR(S):
CORPORATE SOURCE:
SOURCE:
SOURCE:
Dep. Chim. Fac. Sci. Rabat, Morocco
Bulletin de la Societe Chimique de France (1991), (March-April), 255-9
CODEN: BSCFAS; ISSN: 0037-8968
JOURNAI
French
OTHER SOURCE(S):
G1

AB New imidazole (pyrazino and diazepino) benzimidazoles, e.g. I (R = H, Me, n = 1,2,3), II (n = 1,2,3), and III (n = 1,2), were prepared by reaction between 2-aminoalkyl (and aryl) benzimidazoles and dibromoalkanes Br(CH2)nBr (n = 1,2,3) under phase transfer catalysis conditions. These products were characterized by lH-NNR, IR, MS, and microanal.

IT 135075-12-69

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, from o-phenylenediamine, amino acid, and dibromoalkane)
RN 135075-12-6 HCAPLUS
CN 5H-Pyrrolo[1', 2':3, 4]imidazo[1,5-a]benzimidazole, 1,2,3,1lb-tetrahydro-, (S)- (SCI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 35 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

L69 ANSWER 35 OF 63
ACCESSION NUMBER:
DOCUMENT NUMBER:
114:228624 HCAPLUS
114:228624 HCAPLUS
114:228624 HCAPLUS
114:228624 HCAPLUS
114:228624 HCAPLUS
116:228624 HCAP

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
**********				
JP 01272590	A2	19891031	JP 1988-98053	19880422
JP 2568248	B2	19961225		
PRIORITY APPLN. INFO.:			JP 1988-98053	19880422

OTHER SOURCE(S): MARPAT 114:228624

GI For diagram(s), see printed CA Issue.

AB Title compds. I [Y = C,N; R = Q1, R1 = H, protecting group; R2 = H, cyclopropyl-, cyano-, carbamoyl-, or (protected) CO2H-substituted alkyl, Q: R3 = (protected) CO2H, carboxylate; R4 = H, (N-alkyl or N,N-dialkyl)carbamoyl-, cyano-, or (protected) CO2H-substituted alkyl, useful as antibiotics especially for treating drug-resistant bacteria,

(+)-bacteria, and Pseudomonas aeruginosa, are prepared A diastereomer

of I [R = 6,7-dihydro-2-methyl-5H-pyrrolo[1,2-c]imidazolinium-7-yl; R1 = H;R2

Absolute stereochemistry.
Double bond geometry unknown.

(Continued)

127112-07-6 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 2-(2-amino-2-oxoethyl)-7-{[2-

[(diphenylmethoxy)carbonyl]-7-[[(methoxyimino)[2-[(triphenylmethyl)amino]-

4-thiazolyl]acetyl]amino]-5-oxido-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]thio]-6,7-dihydro-, iodide, [6R-(6α,7β)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

ANSWER 35 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) 127112-10-1 HCAPLUS STORM (Continued) 127112-10-1 HCAPLUS H-Pyrrolof(1, 2-c)limidarolium,  $7-[[7-[[(2-\min-4-thiazoly)]][[2-(1,1-\dim-4-thiazoly)]-1-dimethyl-2-oxoethoxy]imino]acetyl]amino]-2-([diphenylmethoxy)carbonyl]-8-oxo-5-thia-1-azabicyclof(4.2.0]oct-2-en-3-yllthio]-6, <math>7$ -dihydro-2-methyl-, iodide,  $[6R-(6\alpha,7\beta)]-(9CI)$  (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

127112-13-4 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 7-[[7-[[[5-{[(1,1-dimethylethoxy)carbonyl]amino]-1,2,4-thiadiazol-3-

yl](ethoxyimino)acetyl]amino]-2-[(diphenylmethoxy)carbonyl]-5-oxido-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, lodide, [6R-(6a.7]p]|- (SCI) (CA INDEX NAME)

Absolute stereochemistry.

127112-14-5 HCAPLUS

L69 ANSWER 35 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN CN 5H-Pyrrolo[1,2-c]imidazolium, 7-[[7-[([5-[[[1,1-dimethylethoxy)carbonyl]amino]-1,2,4-thiadiazol-3-(Continued)

yl] (ethoxyimino)acetyl]amino]-2-[(diphenylmethoxy)carbonyl]-8-oxo-5-thia-l-arabicyclo[4.2:0]oct-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, iodide, [6R-(6a,7B)]- (SGI) (CA INDEX NAME)

Absolute stereochemistry.

• I-

127134-54-7 HCAPLUS
5H-Pyrrolo(1,2-c)imidazolium, 7-[[7-([(2-amino-4-thiazolyl)][[2-{1,1-dimethylethoxy}-1,1-dimethyl-2-oxoethoxy}imino]acetyl]amino]-2-

[(diphenylmethoxy)carbonyl)-5-oxido-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-y1[thio]-6,7-dihydro-2-methyl-, iodide, [6R-(6α,7β)]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

♠ T =

IT 127111-78-8P 127111-79-9P 127111-80-2P

L69 ANSWER 35 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
127111-81-3P 127111-82-4P 127111-85-7P
127111-86-8P 127111-88-1P 127111-90-4P
127111-95-9P 127111-95-9P 127114-33-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study, PREP (Preparation)
(prepn. of, as antibiotic)
RN 12711-178-8 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidarolium, 7-[[7-[[(2-amine-4-thiazoly1)(methoxyimino)acety1]amino]-2-carboxy-8-oxo-5-thia-1-azabioyclo[4,2.0]oct-2-en-3-y1]thio]-6,7-dihydro-2-methy1-, inner salt, [6R-[3(R-1),6a,7B]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

127111-79-9 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 7-{[7-{[(2-amino-4-thiazoly]] (methoxyimino] acety] lamio]-2-carboxy-8-oxo-5-thia-1-azabicyc[64.2.0]oct-2-en-3-y]latio]-6,7-dihydro-2-methyl-, inner salt, [6R-[3(S\*),6a,78]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

L69 ANSWER 35 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

127111-80-2 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 2-(2-amino-2-oxoethyl)-7-[{7-[{{2-amino-4-thiazolyl} (methoxyimino| acetyl | amino| -2-carboxy-8-oxo-5-thia-1-arabicyclo[4.2.0]oct-2-en-3-yllhio]-6, 7-dihydro-, inner salt,  $\{6R-[3(R^*),6\alpha,7\beta]\}$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

127111-81-3 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 2-(2-amino-2-oxoethyl)-7-[[7-[[{2-amino-4-thiazolyl] (methoxyimino| acetyl| amino|-2-carboxy-8-oxo-5-thia-l-azabicyclo[4.2.0] oct-2-en-3-yllhio]-6,7-dihydro-, inner salt, [6R-[3(5\*),6a,78]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

L69 ANSWER 35 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

Absolute stereochemistry.
Double bond geometry unknown.

PAGE 1-A

• c1

● HC1

PAGE 2-A

12711-85-7 HCAPLUS
SH-Pyrrolo[1,2-c|imidazolium, 7-[[7-[([5-amino-1,2,4-thiadiazol-3-y])(ethoxyimino)acety]]amino]-2-carboxy-8-oxo-5-thia-1-arabicyclo[4,2-0]oct-2-en-3-y]lthio]-6,7-dlhydro-2-methyl-, inner salt, [6x-]3[x], 6x,78]]-[9C1] (CA INDEX NAME)

L69 ANSWER 35 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) Absolute stereochemistry.

127111-86-8 HCAPLUS
5H-Pyrrolo[1,2-c|imidazolium, 7-[[7-[[(5-amino-1,2,4-thiadiazol-3-yl)(ethoxyimino)acetyl]emino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4,2.0]oct-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner salt, [6R-[3(5\*),6a,7β]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

127111-89-1 HCAPLUS
5H-Pyrrolo[1,2-c}imidazolium, 7-[[7-{[[5-amino-1,2,4-thiadiazol-3-y1}{[1-carboxy-1-methylethoxy] nino]acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4,2,0]oct-2-en-3-y1]thio]-6,7-dihydro-2-methyl-, inner salt, {6R-[3(R\*),6a,78]}- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 35 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
127111-90-4 HCAPLUS
5H-Pyrrol0[1,2-c|imidazolium, 7-[{7-[{(5-amino-1,2,4-thiadiazol-3-yl){(1-carboxy-1-methylethoxy):mino]acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4,2.0]oct-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner salt,
[6R-[3(3\*),6α,7β]]- (9CI) (CA INDEX NAME)

### Absolute stereochemistry.

127111-95-9 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 7-[[7-[[(5-amino-1,2,4-thiadiazol-3-yl)(hydroxymino)acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl][thio]-6,7-dihydro-2-methyl-, inner salt, [6R-[3(R\*1,6a,78]]- (9CI) (CA INDEX NAME)

### Absolute stereochemistry.

127111-96-0 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 7-[[7-[[(5-amino-1,2,4-thiadiazol-3-yl)(hydroxyimino)acetyl]amino]-2-carboxy-8-oxo-5-thia-1-arabicyc[64.2.0]oct-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner salt, [6R-[3(S\*),6a,7β]]- (9CI) (CA INDEX NAME)

### Absolute stereochemistry.

L69 ANSWER 36 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN

ACCESSION NUMBER: 1990:128873 HCAPLUS

DOCUMENT NUMBER: 112:128873 HCAPLUS

INVENTOR(S): Takeya, Yutaka: Matsurawa, Hiroshi; Iwata, Kaoru

PATENT ASSIGNEE(S): Taijin Ltd., Japan

Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILU ACC. NUM. COUNT: 1

PATENT INFORMATION:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01245084	A2	19890929	JP 1988-72081	19880328
PRIORITY APPLN. INFO.:			JP 1988-72081	19880328

A nonlinear optical material, suited for use in optical switches, memories, and bistable devices, consists of a carbonic acid ester represented by RA(CH:CH)nCH:C(CN)CO2L (R = RIRZM, R3O, R4S, CM, CONRSR6, NRTCOR8, R8; R1-9 = C1-8 hydrocarbyl, H: A = C5-14 aryl; L = C12-25 straight-chain hydrocarbyl; n = 0, 1, 2].

125811-46-1

RET (Reactant); RACT (Reactant or reagent) (reaction of, nonlinear optical material from)

125811-46-3 HCAPLUS

11-Pyrrolo(1,2-c]/midazole, hexahydro-1-(3-methoxyphenyl)-2-phenyl- (9CI) (CA INDEX NAME)

ΙT

L69 ANSWER 35 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

127134-53-6 HCAPLUS SH-Pyrrolo[1,2-c]imidarolium, 7-[{7-{{(2-amino-4-thiazoly1)}{(1-carboxy-1-meth)ethoxy|imino]acety1]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4,2.0]oct-2-en-3-y\_1]thio]-6,7-dihydro-2-methyl-, chloride, monohydrochloride, [6R-{3(S^\*), 6 $\alpha$ , 7 $\beta$ ]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

PAGE 2-A

● c1

L69 ANSWER 37 OF 63
ACCESSION NOMBER:
DOCUMENT NUMBER:
110:75160
Preparation of 6-(1-hydroxyethyl)-2-penem-3carboxylate derivatives as antibacterials
Takemura, Makoto, Zauma, Kunior Nishi, Toshiyuki;
Koda, Hiroko; Sato, Makoto
PATENT ASSIGNEE(S):
DAIICHI SEIVAKU Co., Ltd., Japan
JDN. Kokai Tokkyo Koho, 26 pp.
CODEN: JKXXAF
PAMILY ACC. NUM. COUNT:
1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 63154691	A2	19880627	JP 1986-300810	19861217
JP 2579472	B2	19970205		
PRIORITY APPLN. INFO.:			JP 1986-300810	19861217

AB The title compds. [I; R1 = pyrrolidinoimidazole Q, Q1; R2 = ester residue,
H, or CO2R2 = CO2-; R3 = H, (mono- or di-loweralkyl)carbamoyl, lower alkoxy carbamoyl, morpholinocarbonyl, (un)substituted imidazolyl, (un)substituted thiazolyl, acyl, halo, lower alkyl, lower alkoxy, cycloalkyl, Ph, etc.] were prepared as antibacterials. A solution of 250 mg

(un) substituted thiazolyl, acyl, halo, lower alkyl, lower alkoxy, cycloalkyl, Ph, etc.] were prepared as antibacterials. A solution of 250 mg
p-nitrobenzyl (5R,6S,8R)-2-ethylsulfinyl-6-(1-hydroxyethyl)-2-penem-3-carboxylate in DMF was cooled to -40° and a solution of 440 mg
6,7-dihydro-7-mercapto-5h-pyrrolol(1,2-e]imidazole-CF\$508H in DMF followed by (iso-Pr)2NEt was added. The resulting mixture was stirred at the same temperature for 30 min to give 290 mg (5R,6S,6R)-I [R1 = 6,7-dihydro-5h-pyrrolol1,2-e]imidazol-7-yl, R2 = p-O2NC6HCK12 which was hydrogenolized over Pd/C in THF-phosphate buffer to give an isomeric mixture of (5R,6S,6R)-I (R1 = the same as above, R2 = H). One of the above isomers showed a min. inhibitory concentration of \$0.05 µg/mL against
Staphylococcus aureus.

IT 108308-24-3P 108308-45-9P 108308-41-0P
108308-42-5P 108308-45-9P 108308-41-0P
118776-64-9P 118776-73-1P 118776-75-3P
118776-76-4P 118776-80-0P 118776-61-1P
118776-82-2P 118776-80-0P 118776-61-1P
118776-80-9P 118776-90-2P 118776-91-3P

ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN 118776-92-49 118776-93-59 118776-94-69 118776-97-18776-97-99 118776-98-09 118776-98-07 118777-00-79 118777-01-89 118777-02-99 118859-84-09 118866-55-09 (Continued)

### Absolute stereochemistry.

108308-25-4 HCAPLUS SH-Pyrrolo[1,2-c]imidazolium,  $7-\{[2-carboxy-6-(1-hydroxyethy1)-7-oxo-4-thia-1-axabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methy1-, inner salt, [5R-[3(5*),5<math>\alpha$ ,6 $\alpha$ (R\*)]]- {9CI} (CA INDEX NAME)

### Absolute stereochemistry.

108308-41-4 HCAPLUS  $\frac{1}{2}-\frac{1}{2}$ 

### Absolute stereochemistry.

#### L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

118776-44-6 HCAPLUS 5H-Pyrrolo[1,2-c] imidazolium,  $7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[2-(4-morpholinyl)-2-oxoethyl]-, inner salt, <math>[5R-[3(R^*),5\alpha,6\alpha(R^*)]]-(9CI)$  (CA INDEX NAME)

### Absolute stereochemistry.

118776-73-1 HCAPLUS
SH-Pyrrolo[1,2-climidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[[(1,4,5,6tetrahydro-4-methyl-5,6-dloxo-1,2,4-trizzin-3-yl]thio]methyl]-, inner
salt, [SR-[3[s\*],5a,6(R\*)]]- [951] (CA INDEX NAME)

### Absolute stereochemistry.

l18776-75-3 HCAPLUS SH-Pyrrolo[1,2-climidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azebicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[2-(4-morpholinyl)-2-oxoethyl]-, inner salt,  $[5R-[3(s^*),5a,6a(R^*)]$ -(9CI) (CA INDEX NAME)

# Absolute stereochemistry.

L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

#### Absolute stereochemistry.

108308-46-9 HCAPLUS  $5H-Pyrrolo[1,2-c]imidazolium, 7-[\{2-carboxy-6-\{1-hydroxyethy1\}-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-2-ethyl-6,7-dihydro-, inner salt, <math>[5R-\{3\{R^*\},5\alpha,6\alpha\{R^*\}\}]-$  (9CI) (CA INDEX NAME)

### Absolute stereochemistry.

108308-47-0 RCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-ethyl-6,7-dihydro-, innersalt, [5R-[3(5\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

# Absolute stereochemistry.

#### L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

## Absolute stereochemistry.

118776-77-5 HCAPLUS 5H-Pyrrolo(1,2-c) imidazolium,  $7-[[2-carboxy-6-(1-hydroxyethy1)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[2-(methoxyamino)-2-oxoethy1]-, inner salt, <math>[5R-[3(S^*),5a,6a(R^*)]$ ]- [9CI) (CA INDEX NAME)

### Absolute stereochemistry.

118776-78-6 HCAPLUS 5H-Pyrcolo(1,2-c) inidazolium, 7-([2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-tha-1-azabicyclo(3.2.0) hept-2-en-3-yl)thio]-6, <math>7-dihydro-2-([5-oxo-2-pyrcolidinyl)) methyl]-, inner salt,  $[5R-[3]R^*(S^*)]$ ,  $5\alpha$ ,  $6\alpha$   $(R^*)]$ ]- (9CI) (CA INDEX NAME)

118776-79-7 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 7-[{2-carboxy-6-(1-hydroxyethyl}-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-(2-fluoroethyl)-6,7-dihydro-, inner salt, [5R-[3(R\*),5\omega,6\alpha(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

118776-80-0 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-(2-fluoroethyl)-6,7-dihydro-, inner salt,  $[5R-[3(S^*),5\alpha,6\alpha(R^*)]]-(9CI)$  (CA INDEX NAME)

Absolute stereochemistry.

118776-91-1 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(2-methoxyethyl)-, inner salt, [5R-[3[R\*]],5 $\alpha$ ,6 $\alpha$ (R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

Absolute stereochemistry.

118776-86-6 HCAPLUS 5H-Pyrrolo[1,2-c] imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azobicyclo[3,2-0] hept-2-en-3-yllthio]-6,7-dihydro-2-[(2-pyrimidinylthio)methyl]-, inner salt,  $[5R-[3(R^*),5\alpha,6\alpha(R^*)]]-[9CI)$  (GI NDEX NAME)

Absolute stereochemistry.

118776-87-7 HCAPLUS
5H-Pyrrole[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4thia-1-azabicycle[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[[2pyzimidinylthio]methyl]-, inner salt, [5R-[3[8\*],5u,6u[R\*]]](9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

118776-82-2 HCAPLUS SH-Pyrrolo[1,2-c] imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-y][thio]-6,7-dihydro-2-(2-thienylmethyl)-, inner salt, [5R-[3(R^\*),5 $\alpha$ ,6 $\alpha$ (R^\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

118776-83-3 HCAPLUS SH-Pyrrolo(1,2-climidazolium, 7-{[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azebicyclo(3,2.0)hept-2-en-3-yllthio]-6,7-dihydro-2-(2-thienylmethyl)-, inner salt, [5R-[3(S^+),5 $\alpha$ ,6 $\alpha$ (R^+)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

118776-84-4 HCAPLUS 5H-Pyrrolo[1,2-climidazolium, 7-[[2-carboxy-6-[1-hydroxyethy1]-7-oxo-4-thia-1-azabicyclo[3,2,0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[(methylthio)methyl]-, inner salt, [5R-[3(R\*),5 $\alpha$ ,6 $\alpha$ (R\*)]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

118776-89-9 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[[{1-methyl-1H-tetrazol-5-yl}thio]methyl]-, inner salt, [SR-[3[S\*),5 $\alpha$ ,6 $\alpha$ (R\*)] ]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 118776-90-2 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
2-(2-amino-2-cxoethyl)-6-[(5R,65)-2-carboxy6-[(1R)-1-hydrocxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3yl[thio]-6,7-dihydro-, inner salt (9CI) (CA INDEX NAME)

Absolute stereochemistry.

118776-91-3 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 2-[2-amino-2-oxoethyl)-6-[{2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-arabicyclo[3.2.0]hept-2-en-3-yl]methyl]-, inner salt, [5R-[3(S\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

118776-92-4 HCAPLUS  $\begin{array}{lll} SH-Pyrrolo[1,2-c] imidazolium, & 7-[\{2-carboxy-6-\{1-hydroxyethyl\}-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-y][thio]-6,7-dihydro-2-[2-(methylamino)-2-oxoethyl]-, inner salt, [5R-[3(R^*),5\alpha,6\alpha(R^*)]]- & 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5000 | 1.5$ (CA INDEX NAME)

Absolute stereochemistry.

118776-93-5 HCAPLUS
SH-Pyrclo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2,0]hept-2-en-3-y][thio]-6,7-dihydro-2-[2-(methylamino)-2-oxoethyl)-, inner salt, [5R-[3(5\*),5u,6a(R\*)]]-[9CI] (CA INDEX NAME)

Absolute stereochemistry.

118776-94-6 HCAPLUS 5H-Pyrrolo(1,2-c)imidazolium, 7-{[2-carboxy-6-(1-hydroxyethy1)-7-oxo-4-

thia-1-azabicyclo(3.2.0)hept-2-en-3-yl]thio]-2-[2-(ethylamino)-2-oxoethyl]-6,7-dihydro-, inner salt, [5R-[3(R\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

(Continued) L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

118776-98-0 HCAPLUS 5H-Pyrrole[1,2-c] imidazolium, 2-(3-amino-3-oxopropyl)-7-[[2-carboxy-6-{1-hydroxyethyl}-7-oxo-4-thia-1-azabicyclo[3,2,0]hept-2-en-3-yl]thio]-6,7-dihydro-, inner salt, [SR-[3(R^\*),5a,6a(R^\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

118776-99-1 HCAPLUS 5H-Pyrrolo(1,2-c)imidazolium, 2-(3-amino-3-oxopropyl)-7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo(3,2,0)hept-2-en-3-yl]thio)-6,7-dihydro-, inner salt, [5R-[3(8^\*),5 $\alpha$ ,6 $\alpha$ (R^\*)]]- (9CI) (CA INDEX RN CN

Absolute stereochemistry.

118777-00-7 HCAPLUS SH-Pyrrolo(1,2-c)imidazolium, 7-[(2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo(3.2.01hept-2-en-3-yl]thio]-6,7-dihydro-2-(2-oxo-2-phenylethyl)-, inner salt, [SR-[3(R\*),5a,6a(R\*))]- [9CI] (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

118776-95-7 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-[2-(ethylamino)-2-oxoethyl]-6,7-dihydro-, inner salt, [5R-[3(5\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

118776-96-8 HCAPLUS 5H-Pyrrolo[1,2-c] imidazolium,  $7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2.0] hept-2-en-3-y] thio]-2-[2-(dimethylamino)-2-oxoethyl]-6,7-dihydro-, inner salt, <math>[5R-[3(R^*),5a,6a(R^*)]]-(9CI)$  (CA INDEX NAME)

Absolute stereochemistry.

THE/10-97-9 MCAPLUS SH-Pyrrolo[1,2-c] imidazolium,  $7-[\{2-carboxy-6-\{1-hydroxyethyl\}-7-oxo-4-thia-1-azabicyclo[3,2.0] hept-2-en-3-yl]thio]-2-[2-(dimethylamino)-2-oxoethyl]-6, <math>7$ -dihydro-, inner salt,  $[5R-[3]S^*]$ ,  $5\alpha$ ,  $6\alpha$  (R\*)]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

118777-01-8 HCAPLUS SH-Pyrrolo[1,2~c] imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(2-oxo-2-phenylethyl)-, inner salt, [5R-[3(5\*),5a,6a(R\*)]]- [9CI] (CA INDEX NAME)

Absolute stereochemistry.

thia-1-azabicyclo(3.2.0)hept-2-en-3-yl]thio]-6,7-dihydro-2-[[(1-methyl-1H-tetrazol-5-yl)thio]methyl]-, inner salt, [5R-[3(R\*),5a,6a(R\*)]]- [9C1] (CA INDEX NAME)

Absolute stereochemistry.

118859-84-0 HCAPLUS
SH-Pyrrolo[1,2-cimidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(2-methoxyethyl)-, inner salt, [5R-[3(S\*),5a,6a(R\*)]]- (9CI) (CA INDEX NRME)

Absolute stereochemistry.

Page 249

118776-36-6P 118776-38-8P 118776-54-8P
118776-56-0P 118776-57-1P 118776-53-3P
118776-60-6P 118776-62-8P 118776-63-9P
118776-63-1P 118776-62-2P 118776-68-9P
118776-70-8P 118776-72-0P
RI: SPN (Synthetic preparation); PREP (Preparation)
(preparation of, as intermediate for penem antibacterial)
118776-36-6 NCAPLUS
5H-Pytrolo[1, 2-c]imidazolium, 6,7-dihydro-7-[[(4-methoxyphenyl]methyl]thio]-2-[[(1,4,5,6-tetrahydro-4-methyl-5,6-dioxo-1,2,4-triazin-3-yl)thio]methyl]-, iodide (9CI) (CA INDEX NAME)

RN 118776-56-0 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
6,7-dihydro-7-mercapto-2-{2-(methoxyamino)-2-oxoethyl}-, salt with trifluoromethanesulfonic acid {1:1} (9CI) (CA INDEX NAME)

CH 1

CRN 118776-55-9 CMF C9 H14 N3 O2 S

2 CM

118776-57-1 MCAPLUS
5M-Pyrcolo[1,2-c]imidazolium, 6,7-dihydro-7-[[[4-methoxypheny]]methyl]thio]-2-[[5-oxo-2-pyrcolidiny]]methyl]-, 10dide

(CA INDEX NAME)

• r-

118776-59-3 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6,7-dihydro-7-mercapto-2-[(5-oxo-2-

ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) 118776-38-8 HCAPLUS 5H-Pytrolo[1,2-c]imidazolium, 6,7-dihydro-7-mercapto-2-[[(1,4,5,6-tetrahydro-4-methyl-5,6-dioxo-1,2,4-triazin-3-yl]thio[methyl]-, salt with trifluoromethanesulfonic acid [1:1] [9CI] (CA INDEX NAME)

CH 1

CRN 118776-37-7 CMF C11 H14 N5 O2 S2

CM 2

CRN 37181-39-8 CMF C F3 03 5

118776-54-8 HCAPLUS
5M-Pyrrolo[1,2-c]imidazolium,
-dihydro-2-[2-(methoxyamino]-2-oxoethyl]7-[[(4-methoxyphenyl)methyl]thio]-, bromide (9CI) (CA INDEX NAME)

• Br

L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) pyrrolidinyl)methyl]-, salt with trifluoromethanesulfonic acid (1:1) (9CI)

(CA INDEX NAME)

CM 1

2

CRN 37181-39-8 CMF C F3 03 S

118776-60-6 HCAPLUS
SH-Pyrrolo[1,2-c|imidazolium, 2-(2-fluoroethyl)-6,7-dihydro-7-[[(4-methoxyphenyl methyl]thlo]-, bromide (9CI) (CA INDEX NAME)

● Br

118776-62-8 HCAPLUS
5H-Pyrrolo[1,2-c|imidazolium, 2-(2-fluoroethyl)-6,7-dihydro-7-mercapto-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 118776-61-7 CMF C8 H12 F N2 S

ан 2

CRN 37181-39-8 CMF C F3 03 S

118776-63-9 HCAPLUS
5H-Pyrrolo(1,2-c)imidazolium, 6,7-dihydro-7-{{{4-methoxyphenyl}methyl}thio}-2-{2-thiazolylmethyl}-, chloride (9CI) (CA INDEX NAME)

• c1-

RN 118776-65-1 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
6,7-dihydro-7-mercapto-2-(2-thiazolylmethyl), salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 118776-64-0 CMF C10 H12 N3 52

L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

CM

118776-70-8 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6,7-dihydro-7-[[{4-methoxyphenyl]methyl]thio]-2-[{2-pyrimidinylthio}methyl}-, iodide (9CI)
(CA INDEX NAME)

• T-

118776-72-0 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6,7-dihydro-7-mercapto-2-{(2-pyrimidinylthio)methyl-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 118776-71-9 CMF C11 H13 N4 S2

L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

CH 2

CRN 37181-39-8 CMF C F3 03 S

118776-66-2 HCAPLUS
5H-Pyrrolo[1,2-c]imidezolium, 6,7-dihydro-7-{{{4-methoxyphenyl}methyl}thio]-2-{{methylthio}methyl}-, chloride (9CI) (CA INDEX NAME)

• c1-

118776-68-4 HCAPLUS
5H-Pyrrolo[1,2-c|imidazolium, 6,7-dihydro-7-mercapto-2[(methylthio)methyl]-, salt with trifluoromethanesulfonic acid (1:1) (9CI)

(CA INDEX NAME)

CM 1

CRN 118776-67-3 CMF C8 H13 N2 S2

L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

CM 2

118776-45-7P 118776-47-9P 118776-48-0P 118776-50-4P 118776-52-6P 118776-90-9P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as intermediate for penem antibacterials) 118776-63-7 HCAPLUS SH-Pyrrolo[1, 2-c]imidazolium, 6,7-dihydro-7-[[(4-methoxyphenyl)methyl]thio]-2-[2-(methylamino)-2-oxoethyl]-, bromide (9CI) (CA INDEX NAME)

● Br-

RN 118776-47-9 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
6,7-dihydro-7-mercapto-2-[2-(methylamino)-2oxoethyl]-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAMES

CH 1

CRN 118776-46-8 CMF C9 H14 N3 O S

CM 2

CRN 37181-39-8 CMF C F3 O3 S

P- C- 503

118776-48-0 HCAPLUS 5H-Pyrrolo[1,2-c|imidazolium, 2-(2-amino-2-oxoethyl)-6,7-dihydro-7-[[(4-methoxyphenyl)methyl]thio]-, chloride (SCI) (CA INDEX NAME)

• c1 -

118776-50-4 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 2-(2-amino-2-oxoethyl)-6,7-dihydro-7-mercapto-, salt with trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 118776-49-1 CMF C8 H12 N3 O S

L69 ANSMER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
CN 5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-[[[1,4,5,6tetrahydro-4-methyl-5,6-dioxo-1,2,4-triazin-3-yl]thio]methyl]-, inner
salt, [5R-[3(R\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 37 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

CM 2

CRN 37181-39-8 CMF C F3 03 S

118776-52-6 HCAPLUS
SH-Pyrrolof1,2-c]imidazolium, 6,7-dihydro-7-mercapto-2-methyl-, salt with
trifluoromethanesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 118776-51-5 CMF C7 H11 N2 S

2

CRN 37181-39-8 CMF C F3 03 S

118776-88-8 HCAPLUS

L69 ANSWER 38 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1988:59768 HCAPLUS
DOCUMENT NUMBER: 108:59768
TITLE: Late Quaternary Mediterranean sapropels. II.

Organic

geochemistry and palynology of S1 sapropels and associated sediments

AUTHOR(S): Ten Haven, H. L.; Baas, M.; De Leeuw, J. W.; Schenck, P. A.; Brinkhuis, H.

CORPORATE SOURCE: Dep. Chem. Chem. Eng., Delft Univ. Technol., Delft, 2628 R2, Neth.

SOURCE: Chemical Geology (1987), 64(1-2), 149-67

CODDENT TYPE: Journal

LANGUAGE: English

AB The organic matter of S1 sapropels is of a mixed marine, terrigenous, and bacterial origin. A trend of relatively increasing amts. of continent-derived organic matter towards more seaward and deeper realms can

be observed from both palynol, and organic geochem. data. This trend is supported to some extent by \$0.33C-values of the organic matter. The sapropelic intervals deposited on the Nile Cone are characterized by expanded thicknesses and a diluted organic C content because of a higher sedimentation rate. The environmental conditions (in terms of preservation) during sapropel formation over the eastern Mediterranean were probably not uniform. At site 29, the conditions were favorable for the deposition of sapropel with a higher organic C content than at the

locations. This might have been caused by better preservation

locations. This might have been caused by better preservation conditions.

Increasing discharge from the Nile River was the driving force for formation of the S1 sapropels. Based on this assumption a model for sapropel formation is proposed.

IT 236-71-5

236-1-5
RL: GOC (Geological or astronomical occurrence); OCCU (Occurrence) (in sapropels, of late Quaternary, of eastern Mediterranean) 236-11-5 HCAPLUS
SH-Imidazo[5,1-b:4,3-b']bisthiazole (8CI, 9CI) (CA INDEX NAME)

M

ANSWER 39 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of) 112565-43-2 HCAPLUS SH-Pytrolo[1',2':3,4]imidazo[1,5-a]indole, 1,2,3,1lb-tetrahydro-11-(methoxymethyl)- (9C1) (CA INDEX NAME)

L69 ANSWER 39 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1988:55922 HCAPLUS
108:55922 Synthesis of some hexahydroazocino{4,3-b}indoles, a
tetra- and two
hexahydropyrrolo{1',2':1,2jpyrrolo{3,4b}indoles, and a
tetrahydropyrrolo{2',1':5,1jmidazo{3}
,4-ajindole. Crystal structure determination of
1,2,3,4-tetrahydro-2-(phenoxycarbonyl)-7(phenylsulfonyl)azocino{4,3-b}indol-6(5H)-one
AUTHOR(S):
Street, Jonathan D.: Harris, Martin; Blahop, David

Heatley, Frank: Beddoes, Roy L.: Mills, Owen S.: Joule, John A. Chem. Dep., Univ. Manchester, Manchester, M13 9PL, UK Journal of the Chemical Society, Perkin Transactions 1: Organic and Bio-Organic Chemistry (1972-1999) (1987), (7), 1599-606 CODEM: JCPRB4: ISSN: 0300-922X Journal CORPORATE SOURCE: SOURCE:

DOCUMENT TYPE:

LANGUAGE: OTHER SOURCE(S): GI English CASREACT 108:55922

Azocinoindoles I (R = H, SO2Ph; R1 = H, R2 = OH; R1R2 = O; R3 = CH2Ph) were prepared from 1-phenylsulfonylindole by introducing the appropriate side chain at C-2 via lithiation, and then intramol. Nannich cyclization. I (R = SO2Ph, R1R2 = O, R = CO2Ph, II) was prepared from I (R = SO2Ph, R1R2

= 0, R3 = CH2Ph, III). The mol. structure of II was determined by x-ray crystal structure anal. The benzyl group of II was also replaced by other

urethane groups. Cleavage of the urethanes gave pyrrolopyrroloindoles, e.g., IV. Reaction of 2-indol-2-ylpyrrolidine with CH20 in methanolic methoxide gave pyrroloimidazoindole V.

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

Penem derivs. I [R1 = H, alkyl, hydroxyalkyl; CO2R2 = CO2H, CO2-; R2 = ester group, protective group; R3 = {un}substituted bicycloheterocyclyl} and their salts, useful as antibacterial agents with extremely wide antibacterial spectrum, were prepared by reacting 2-substituted sulfinyl derivs. Of penem with HSR3 and then optionally removing protective group(s) and further alkylating the reaction product or vice versa. Et urocanate-HCl was alkalinated and reacted with 4-MeOCSHGHZSH and the product Et 3-{inidazol-4-yl}-3-[c-methoxybenzylthio]propionate was converted in 4 steps to pyrroloimidazole salt II. This reacted with p-nitrobenzyl (5R,65,6R)-2-ethylsulfinyl-6-[1-hydroxyethyl]-2-penem-3-carboxylate to give the sulfide (5R,65,6R)-III (R2 = 4-O2NC6H4CH2), hydrogenolysis of which over 101 Pd/C gave (5R,65,6R)-III (R = H) (IV) as isomers A and B. The min. inhibitory concentration of isomer A of IV nat E. isomers A and B. The min. inhibitory concentration of isomer A of IV against E.

coli NIHJ was 0.1 µg/mL, whereas that of carbamate V was 0.39 µg/mL.

108308-48-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and deprotection of)

RN 108308-48-1 HCRPLUS

CN 5H-Pyrrolc1, 2-c1 imidazolium,
2-ethyl-6,7-dihydro-7-[[6-(1-hydroxyethyl)-2-

[{(4-nitrophenyl)methoxy}carbonyl}-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl}thio]-, iodide, [5R-[5α, 6α(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1987:213640 HCAPLUS
TITLE: 106:213640 HCAPLUS
INVENTOR(5): 840 HCAPLUS
ANALOGY ARE SERVICE STREET ASSIGNEE(S): 50URCE: 106:213640 HCAPLUS
DOCUMENT TYPE: PALENT ASPIL, 180 pp.
CODEN: EXXXDW
DOCUMENT TYPE: PALENT ASPIL, 180 pp.

English

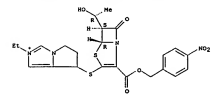
FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.		DATE	APPLICATION NO.	DATE
	A2	19870204		19860617
		19870701		
EP 210883	B1	19950201		
		, GB, IT,	LI, NL, SE	
CA 1339860	A1		CA 1986-511536	
ZA 8604472	A	19870225	ZA 1986-4472	19860616
DK 8602835	A	19861218	DK 1986-2835 FI 1986-2566	19860617
DK 171644	B1	19970303		
FI 8602566	A	19861218	FI 1986-2566	19860617
21 0/410		19920031		
FI 87218	С	19921210		
NO 8602411 NO 167982 NO 167982	A	19861218	NO 1986-2411	19860617
NO 167982	В	19910923		
NO 167982	С	19920102		
AU 8658909	A1	19861224	AU 1986-58909	19860617
AU 593558	B2	19900215		
JP 62149683	A2	19870703	JP 1986-141171	19860617
JP 08026040	B4	19960313		
ES 556137 US 4962202 AU 8944784	A1	19880101	ES 1986-556137	19860617
US 4962202	A	19901009	US 1988-117617	19880111
AU 8944784	A1	19900308	AU 1989-44784	19891117
AU 626606	B2	19920806		
US 5079357	A	19920107		19891207
JP 06184146	A2	19940705		19930617
PRIORITY APPLN. INFO.:			JP 1985-131394 A	19850617
			JP 1985-213420 A	19850926
			US 1986-875228 B	2 19860617
			US 1987-38640 B	3 19870415

OTHER SOURCE(S): MARPAT 106:213640

(Continued)

06/28/2006



• I-

108308-26-5P
RL: RCT (Reactant): SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and reduction of)
108308-26-5 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6,7-dihydro-7-[[6-(1-hydroxyethyl)-2-[[4-

nitrophenyl)methoxy|carbonyl}-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-methyl-, iodide, [5R-[5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-24-3P 108308-25-4P 108308-27-6P 108308-28-7P 108308-29-8P 108308-30-1P 108308-31-2P 108308-32-3P 108308-33-4P

• r-

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

108308-28-7 HCAPLUS
5H-Pyrrolo(1,2-c)imidazolium, 5-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(2-oxopropyl)-, inner salt, [5R-[3(5\*),5α,6α(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-29-8 HCAPLUS 5H-Pyrrolof[1,2-c]imidazolium,  $7-[\{2-carboxy-6-(1-hydroxyethy1)-7-oxo-4-thia-1-azabicyclof[3,2.0]$ hept-2-en-3-yl[thio]-2-(cyclopropylcarbony1)-6,7-dihydro-, inner salt,  $[5R-[3(s^*),5\alpha,6\alpha(R^*)]]-(9CI)$  (CA INDEX NAME)

Absolute stereochemistry.

RN 108308-30-1 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
2-(cyclopropylcarbonyl)-6,7-dihydro-7-[[6-[1hydroxyethyl)-2-[[(4-nitrophenyl]methoxy]carbonyl]-7-oxo-4-thia-1azabicyclo[3.2.0]hept-2-en-3-yl]thio]-, lodide, [5R[5a, 6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
108308-34-59 108308-35-69 108308-35-79
108308-37-89 108308-38-99 108308-39-09
108308-40-39 108308-41-79 108308-42-59
108308-46-99 108308-41-79 108308-42-59
108308-46-99 108308-47-79 108308-92-9
108308-50-59 108308-31-69 108308-52-79
108308-50-59 108308-31-69 108308-51-59
108325-52-69
RL: BAC (Biological activity or effector, except adverse); BSU
(Biological study); PREP (Preparation)
(prepn. of, as antibacterial)
RN 108308-24-3 HCAPLUS
CN 5H-Pyrrolo[1,2-c]midscolium, 7-{[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner salt, [5R-(3(R1),50,60(R1)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-25-4 HCAPLUS
5H-Pyrrolo[1, 2-c]imidazolium, 7-[{2-carboxy-6-(1-hydroxyethyl)-7-oxo-4thia-1-azabicyclo[3,2,0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner
salt, [5R-[3(5\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-27-6 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 5-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(2-oxopropyl)-, inner salt, [5R-[3(R\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

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108308-31-2 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-(cyanomethyl)-6,7-dihydro-, inner salt,  $[5R-[3(R^*),5\alpha,6\alpha(R^*)]]-(9CI)$  (CA INDEX NAME)

Absolute stereochemistry.

108308-32-3 HCAPLUS 5H-Pyrrolo(1,2-c)imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

-1-azabicyclo[3.2.0]hept-2-en-3-y1]thio]-2-(cyanomethyl)-6,7-dihydro-, inner salt, [5R-[3(S\*),5a,6a(R\*)]}- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-33-4 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 7-([2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

Searched by Jason M. Nolan Page 254

Absolute stereochemistry

108308-34-5 HCAPLUS
5H-Pyrrolo(1,2-c)imidazolium, 7-[{2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-(carboxymethyl)-6,7-dihydro, inner salt, monosodium salt, [ $SR-[3(S^*),5\alpha,6\alpha(R^*)]$ ]- (9C1) (CA INDEX NAME)

Absolute stereochemistry.

● Na

108308-35-6 HCAPLUS  $7-\{[2-carboxy-6-(1-hydroxyethy1)-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dlhydro-2-(2-propeny1)-, inner salt, <math>[5R-[3(R^*),5\alpha,6\alpha(R^*)]]-\{9CI\}$  (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

108308-39-0 HCAPLUS 5H-Pyrrolo(1,2-c]imidazolium, 7-{[2-carboxy-6-{1-hydroxyethyl}-7-oxo-4-thia-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(2-methoxy-2-oxoethyl)-, inner salt, [5R-[3(R\*),5 $\alpha$ ,6 $\alpha$ (R\*)]}- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-40-3 HCAPLUS 5H-Pyrrolo(1,2-c]imidazolium, 6,7-dihydro-7-[[6-(1-hydroxyethyl)-2-([(4-

nitrophenyl)methoxylcarbonyl]-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio}-2-(2-methoxy-2-oxoethyl)-, bromide, [5R-[5 $\alpha$ , 6 $\alpha$ (R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● Br

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

06/28/2006

108308-36-7 HCAPLUS  $5H-Pyrrolo[1,2-c]imidazolium, 7-{{2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(2-propenyl)-, inner salt, <math>(5R-{3(S^*)}, 5\alpha, 6\alpha(R^*))$ - (QCI INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

Absolute stereochemistry.

ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-, inner salt, [SR-[3(R\*),5a,6a(R\*)]]- (SCI) (CA INDEX NAME)

Absolute stereochemistry.

108308-42-5 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 2-(2-amino-2-oxoethyl)-7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-, inner salt, [5R-[3(S\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 108308-43-6 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
2-(2-amino-2-oxoethyl)-6,7-dihydro-7-[[6-(1-hydroxyethyl)-2-[[(4-nitrophenyl)methoxy]carbonyl]-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl[thio]-, iodide, [5R-[5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Searched by Jason M. Nolan

• r-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(phenylmethyl)-, inner salt, [5R-[3(R^\*),5 $\alpha$ ,6 $\alpha$ (R\*)]}- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-45-8 HCAPLUS 5H-Pyrrolo(1,2-c)imidazolium, 7-[(2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(phenylmethyl)-, inner salt, [5R-[3( $S^*$ ),5 $\alpha$ ,6 $\alpha$ (R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) CN 5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(4-methoxy-2,4-dioxobutyl)-, inner salt, [5R-[3(5\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-51-6 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium,  $7-[\{2-carboxy-6-\{1-hydroxyethyl\}-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl\}thio]-2-[\{4-fluorophenyl\}methyl\}-6,7-dihydro-, inner salt, <math>\{5R-\{3(R^*),5\alpha,6\alpha(R^*)\}\}-\{9CI\}$  (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

108309-32-6 HCAPLUS
5H-Pyrrolo[1, 2-c]imidazolium, 6-{{2-carboxy-6-{1-hydroxyethyl}}-7-oxo-4-thia-1-azabicyclo[3,2,0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner salt, [5R-[3(R\*),5d,6d(R\*)]]- (9CI) (CA INDEX NAME)

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

108308-46-9 HCAPLUS
5H-Pyrrolo[1, 2-c]imidazolium, 7-{[2-carboxy-6-{1-hydroxyethyl}-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-ethyl-6,7-dihydro-, inner salt, {5R-[3(R\*),5a,6a(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-47-0 HCAPLUS 5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-2-ethyl-6,7-dihydro-, inner salt, [5R-[3(5\*),5 $\alpha$ ,6 $\alpha$ (R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-49-2 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-

thia-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-(4-methoxy-2,4-dioxobutyl)-, inner salt, [5R-[3[R\*],5a,6a[R\*]]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

108308-50-5 HCAPLUS

L69 ANSWER 40 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN Absolute stereochemistry. (Continued)

108325-51-5 HCAPLUS  $\begin{array}{lll} & & & \\$ 

Absolute stereochemistry.

108325-52-6 HCAPLUS 5H-Pyrrolo[1,2-c] imidazolium,  $7-\{[2-carboxy-6-(1-hydroxyethyl)-7-oxo-4-thia-1-azabicyclo[3.2.0]$  hept-2-en-3-yllthio]-6,7-dihydro-2-(2-methoxy-2-oxoethyl)-, inner salt,  $[5R-\{3(s^*),5a,6a(R^*)\}]-$  (9CI) (CA INDEX NAME)

Absolute stereochemistry.

K

L69 ANSWER 41 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

104285-15-6 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 5-[[2-carboxy-6-(1-hydroxyethyl)-7-oxo-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner salt (9CI) (CA INDEX NAME)

104285-16-7 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 6-[(2-carboxy-6-(1-hydroxyethyl)-7-oxo-1-azabicyclo[3.2.0]hept-2-en-3-yl}thio]-6,7-dihydro-2-mathyl-, inner salt (9C1) (CA INDEX NAME)

104285-17-8 HCAPLUS
5H-Pyrrolo[1,2-c]imidazolium, 7-[[2-carboxy-6-(1-hydroxyethyl]-7-oxo-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner salt (SCI) (CA INDEX NAME)

L69 ANSWER 41 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1987:67000 HCAPLUS DOCUMENT NUMBER: 106:67000 HCAPLUS 106:67000 HCAPLUS Cacapenems having a 2-minor of Cacapenems Carbapenems having a 2-quaternary hetero-arvlalkylthio substituent
Christensen, Burton G.; Johnston, David B. R.;
Schmitt, Susan M.
Merck and Co., Inc., USA
Eur. Pat. Appl., 194 pp.
CODEN: EPXXDW
Patent INVENTOR (S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
EP 169410	A1 19860	129 EP 1985-108133	19850701
EP 169410	B1 19930	224	
R: AT, BE, CH,	DE, FR, GB,	IT, LI, LU, NL, SE	
CA 1285940	A1 19910	709 CA 1985-486090	19850628
DK 8502974	A 19860	314 DK 1985-2974	19850701
ES 544753	A1 19860	916 ES 1985-544753	19850701
AT 85978	E 19930	315 AT 1985-108133	19850701
JP 61083184	A2 19860	426 JP 1985-145660	19850702
PRIORITY APPLN. INFO.:		US 1984-626580 A	19840702
		EP 1985-108133 A	19850701

For diagram(s), see printed CA Issue. The title compds. I [L = (un)substituted Cl-6 alkyl, C2-6 alkenyl, C3-6 cycloalkyl, etc.; X completes an (un)substituted mono- or bicyclic heterocyclyl: Y = CO2H, CO2R; R = removable CO2H protecting group, CO2M;

= alkali metal), their esters and salts, useful as antibiotics (no data)
were prepared Thus, p-nitrobenzyl (5R,6S)-2-{(diphenylphosphono)oxyl-6{|(R|-hydroxyethyl|carbapen-2-em-3-carboxylate in MeCN was treated with
1-(2-mercaptoethyl)pyridinium nitrate in DMSO and with EtN(CHMe2)2 to

give  $(5R,6S)-6-\{1\{R\}-hydroxyethyl\}-2-[(2-pyridinioethyl)thio] carbapen-2-em-3-(5R,6S)-6-\{1\{R\}-hydroxyethyl\}-2-[(2-pyridinioethyl)thio] carbapen-2-em-3-(5R,6S)-6-\{1\{R\}-hydroxyethyl\}-2-[(2-pyridinioethyl)thio] carbapen-2-em-3-(5R,6S)-6-\{1\{R\}-hydroxyethyl\}-2-[(2-pyridinioethyl)thio] carbapen-2-em-3-(5R,6S)-6-(5$ 

Carboxylate. 104262-91-19 104285-15-69 104285-16-79 104285-17-89 IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological

(Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (preparation of, as antibiotic)

RN 104262-91-1 HCAPLUS
CN 5H-Pytrolo(1,2-c]imidazolium,
2-[2-[[2-carboxy-6-(1-hydroxyethy1)-7-oxo-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]ethyl]-6,7-dihydro-, inner salt, [5R-[5\alpha, 6\alpha(R^\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 41 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

L69 ANSWER 42 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1987:66999 HCAPLUS
DOCUMENT NUMBER: 106:66999
1-Methylcarbapenems having a 2-quaternary heteroarylalkylthio substituent
Christensen, Burton G.; Johnston, David B. R.;
Schmitt, Susan M.
Herck and Co., Inc., USA
SOURCE: EVER AND COUNT: 1000: EPXXDW
DOCUMENT TYPE: EPXID PATENT
LANGUAGE: EPXID PATENT
ENGLISH TORSHATION: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA	TENT NO.		KIND	DATE	APPLICATION NO.	DATE
EP	168707		A1	19860122	EP 1985-108134	19850701
	R: AT,	BE, CH,	DE, FR	, GB, IT,	LI, LU, NL, SE	
ÇA	1273013		Al	19900821	CA 1985-486070	19850628
DK	8502976		Α	19860103	DK 1985-2976	19850701
ES	544754		A1	19860916	ES 1985-544754	19850701
JP	61063679	•	A2	19860401	JP 1985-145661	19850702
PRIORIT	Y APPLN.	INFO.:			US 1984-626822 F	19840702

For diagram(s), see printed CA Issue.
The title compds. I (L = (un)substituted Cl-6 alkyl, C2-6 alkenyl, C3-6 cycloalkyl, etc.; X completes an (un)substituted mono- or bicyclic heterocycle; Y = CO2H, CO2H; R = removable protecting group, CO2H; M = alkali metal), their esters and salts, useful as antibiotics (no data), were prepared Thus, p-nitrobenzyl (55,65)-2-[(diphenylphosphono)oxy]-6-[1(R)-hydroxyethyl]-[R]-methylcarbapen-2-em-3-carboxylate and 3-hydroxy-1-(mercaptoethyl)pyridinium nitrate in MeCONMe2 was treated

EtNPr2-iso2, then diluted with BuOH, EtOAc, and H2O, the pH adjusted with N-methylmorpholine-HCl, treated with Pd(OH)2/C and hydrogenated to give

Na

(5S,6S)-6-[1(R)-hydroxyethyl]-1(R)-methyl-2-[2-(3-oxidopyridinium) ethylthiolcarbapen-2-em-3-carboxylate.

IT 10391-125-7P 10391-26-8P 103911-27-9P

103965-56-6P

RI: BAC (Biological activity or effector, except adverse); BSU

(Biological

study unclassified); SPN (Synthetic preparation); BIOL (Biological)

(Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (preparation of, as antibiotic) (preparation of, as antibiotic) (preparation of, as antibiotic) (preparation) (preparat

L69 ANSWER 42 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 103911-26-8 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
6-[27-carboxy-6-(1-hydroxyethyl)-4-methyl-7oxo-1-azabicyclo[3.2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner
sait (9C1) (CA INDEX NAME)

RN 103911-27-9 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
7-[[2-carboxy-6-(1-hydroxyethyl)-4-methyl-7oxo-1-azabicyclo[3,2.0]hept-2-en-3-yl]thio]-6,7-dihydro-2-methyl-, inner
salt (9CI) (CA INDEX NAME)

RN 103965-56-6 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazolium,
2-[2-[[2-carboxy-6-(1-hydroxyethyl)-4-methyl7-oxo-1-azabicy-lo[3,2.0]hept-2-en-3-yl]thio]ethyl]-6,7-dihydro-, inner
salt, [4R-[4α,5β,6β(R\*)]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



L69 ANSWER 42 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

Substituted C2 elent

L69 ANSWER 43 OF 63 RCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:
DOCUMENT NUMBER:
1986:5820 HCAPLUS
104:5820
Lithium aluminum hydride reduction with a ring
closure. A facile synthesis of 1,3dialkylimidazolidines and 1,3-disubstituted
4-imidazolines from 4-amino acid derivatives
Kiyooka, Syunichi; Goto, Fumitaka; Fujiyama, Ryoji;
Suzuki, Kojiro
CORPORATE SOURCE: Fac. Sci., Kochi Univ., Kochi, 780, Japan
Kochi Daigaku Rigakubu Kiyo, Kagaku (1985), 6, 15-20
CODEN: KDRKDD; ISSN: 0389-0279
JOURNAL
GI

DOCUMENT TYPE: LANGUAGE: GI

N-Isopropyl-N-(benzyloxycarbonyl)prolinamide was treated with LiAlH4 in THF to give 3-isopropyl-1,3-diazabicyclo(3.3.0)octane (I; R = Me2CH).

Similarly, N-isopropyl-N-(benzyloxycarbonyl)sarcosinamide and N-phenyl-N-(benzyloxycarbonyl)prolinamide gave 1-isopropyl-3-methylimidazolidine and 3-phenyl-1,3-diazabicyclo(3.3.0)octane (I; R = Ph), resp.

SPMO3-62-65 99405-64-8P
RL: SRN (Synthetic preparation); PREP (Preparation)
(preparation of)
99405-62-6 HCAPLUS
IH-Pyrrolo(1,2-c)imidazole, hexahydro-2-(1-methylethyl)- (9CI) (CA INDEX NAME)

99405-64-8 HCAPLUS
1H-Pyrrolo[1,2-c]imidazole, hexahydro-2-phenyl- (9CI) (CA INDEX NAME)



L69 ANSWER 44 OF 63
ACCESSION NUMBER:
DOCUMENT NUMBER:
1985:78770 HCAPLUS
102:78770 BENTIMIDED:
XIII. Reactions of
2-(a-chloro-3',4',5',6'cetrahydrobenzyl)benzimidazole with phenols and
aromatic amines
Saczewski, Franciszek; Foks, Henryk; Sawlewicz, Jozef
CORPORATE SOURCE:
ACTA BOOLOGY
SOURCE:
ACTA BOOLOGY
ACTA BO

CASREACT 102:78770 OTHER SOURCE(S):

111

AB The title benzimidazole derivative (I, R = CI) when around the C6H4CL-4.

C6H4CNL-4) in MeOH gave I (R = OR1). A similar reaction with 4-R2C6H4NH2 (R2 = H, Cl, Me) in the presence of Et3N yielded I (R = NHC6H4R2) (II).

I.HCl (R = Cl) with Et3N in DMF yielded the dimer III. A 1,3-dipole was the intermediate in the investigated reactions. Cyclocondensation of II with CH2O in EtCH gave the corresponding imidazobenzimidazoles IV.

11 9460-98-09-09-0460-90-19 9460-91-29 RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

L69 ANSWER 44 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN CN 1H-Imidazo[1,5-a]benzimidazole,
3-(1-cyclohexen-1-yl)-2,3-dihydro-2-phenyl(9c1) (CA INDEX NAME) (Continued)

94640-90-1 HCAPLUS
1H-Imidazo[1,5-a]benzimidazole, 2-(4-chlorophenyl)-3-(1-cyclohexen-1-yl)2,3-dihydro- (9CI) (CA INDEX NAME)

94640-91-2 HCAPLUS
1H-Tmidazo[1,5-a]benzimidazole, 3-(1-cyclohexen-1-y1)-2,3-dihydro-2-(4-methylphenyl)- (9CI) (CA INDEX NAME)

L69 ANSWER 45 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1984:103247 HCAPLUS
DOCUMENT NUMBER: 100:103247 HCAPLUS
101:03247 HCAPLUS
100:103247 HCAPLUS
200:103247 HCAPLUS

DOCUMENT TYPE: LANGUAGE:

CODEN: JCPRB4; ISSN Journal English CASREACT 100:103247 OTHER SOURCE(S):

N-(Dialkylaminomethyl) succinimides and -glutarimides cyclized to 1,3-diazabicyclo[3.3.0]octanes and -[4.3.0]nonanes, resp., on

irradiation in
MeCN. E.g., irradiation of succinimide I in MeCN gave 26 and 20% yields

the 2 diastereoisomers of diazabicyclooctanes II. N-(Dialkylaminoethyl) aliphatic imides gave azepine- or azocinediones on irradiation, whereas N-(dialkylaminopropyl) derivs. cyclized to give products with a novel perhydro-1,4-diazepine ring. Cyclization of N-(dialkylaminoethyl)maleimide and the analogous 3,4,5,6-tetrahydrophthalimide gave compds. containing a new piperazine ring. 89003-45-2P 89003-46-3P 89003-47-4P 89003-45-2P 89003-49-6P 89003-50-9P RD: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

(preparation of) 89003-45-2 HCAPLUS 5H-Pycrolo(1,2-c)lmidazol-5-one, hexahydro-7a-hydroxy-2-methyl- (9CI)

INDEX NAME)

L69 ANSWER 45 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

89003-46-3 HCAPLUS
SH-Pyrrolo[1,2-c]imidazol-5-one, 1-ethenylhexahydro-7a-hydroxy-2-(2-propenyl)-, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.

89003-47-4 HCAPLUS

5H-Pyrrolo(1,2-c)imidazol-5-one, 1-ethenylhexahydro-7a-hydroxy-2-(2-propenyl)-, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

89003-48-5 HCAPLUS

5H-Pyrrolo[1,2-c]imidazol-5-one, hexahydro-7a-hydroxy-2-methyl-7-phenyl-, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

89003-49-6 HCAPLUS 5H-Pyrrolo[1,2-c]imidazol-5-one, hexahydro-7a-hydroxy-2-methyl-7-phenyl-,

L69 ANSWER 45 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN cis- (9CI) (CA INDEX NAME) (Continued)

Relative stereochemistry.

89003-50-9 HCAPLUS
SH-Pyrcolo[1,2-c|imidazol-5-one, hexahydro-7a-hydroxy-2-methyl-6-phenyl-, trans- (9C1) (CA INDEX NAME) RN CN

Relative stereochemistry.

89003-51-0 HCAPLUS
5H-Pyrrolo[1,2-c]imidazol-5-one, hexahydro-7a-hydroxy-2-methyl-6-phenyl-, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.

83095-08-3P 83095-09-4P IT 83095-08-19 83095-09-4P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of, by photocyclization of
(dialkylaminomethyl) succinimide)
RN 83095-08-3 HCAPLUS
CN 5H-Pyrrolo[1,2-c]imidazol-5-one, 2-ethylhexahydro-7a-hydroxy-1-methyl-,
cis- (9CI) (CA INDEX NAME)

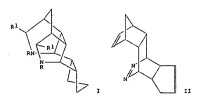
Relative stereochemistry.

L69 ANSWER 46 OF 63
ACCESSION NUMBER:
DOCUMENT NUMBER:
1984:34517 HCAPLUS
100:34517
Preparation of some polycyclic diamine derivatives
Nelsen, Stephen F.; Willi, Mark R.
S. M. McElvain Lab. Org. Chem., Univ. Wisconsin,
Madison, WI, 53706, USA
DOCUMENT TYPE:
LANGUAGE:
DOTHER SOURCE(S):
GI

HCAPLUS COPYRIGHT 2006 ACS on STN
1984:34517 HCAPLUS
100:34517
Preparation of some polycyclic diamine derivatives
Nelsen, Stephen F.; Willi, Mark R.
S. M. McElvain Lab. Org. Chem., Univ. Wisconsin,
Madison, WI, 53706, USA
JOURNAL
LANGUAGE:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
GI

CASREACT 100:34517

OTHER SOURCE(S):



Diezahexacyclohexadecanes I (R = H, Cl, Me, NO, NH2, NMe2, NEL2; R12 = bond) and the related pentacyclic compds. lacking the 2,12 C-C bond were prepared from cyclopentadiene and 2,5-dimethoxy-2,5-dihydrofuran via photolysis of II. Mol. mechanics calcns. on I (R = Me, R12 = bond, R1 = H) are discussed. 87901-37-4P
RL: RCT (Reactant): SPN (Synthetic preparation): PREP (Preparation); RACT (Reactant or reagent) (preparation and reduction of) 87801-37-4 HCAPLUS 2H-1,5,2,4-(Methanonitrilometheno)dicyclopent[cd,q]indolium, dodecahydro-1-methyl-, (la.2a,2aß,4a,4aß,5.alp ha.5aß,8aß,bß,6ß,5s\*)-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME) IT

CM 1

CRN 87801-36-3 CMF C16 H23 N2

L69 ANSWER 45 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

83095-09-4 HCAPLUS
5H-Pyrrolo[1,2-c]imidazol-5-one, 2-ethylhexahydro-7a-hydroxy-1-methyl-,
trana- (9CI) (CA INDEX NAME)

Relative stereochemistry.

L69 ANSWER 46 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

CRN 14874-70-5 CMF B F4 CCI CCS



L69 ANSWER 47 OF 63
ACCESSION NUMBER:
D92:544818 HCAPLUS
TITLE:
AUTHOR(S):
CORPORATE SOURCE:
SOURCE:
CORPORATE SOURCE:
SOURCE:
CORPORATE SOURCE:
CORPORATE SOURCE:
SOURCE:
SOURCE:
CORPORATE SOURCE:
SOURCE:
SOURCE:
CORPORATE SOURCE:
SOURCE:
SOURCE:
CORPORATE SOURCE:
S

CODEN: JRPSDC: ISSN: 0308-2342 DOCUMENT TYPE:

LANGUAGE: OTHER SOURCE(S):

Journal English CASREACT 97:144818

Photochem. cyclization of N-substituted succinimides I [n=1, X=CH20, CH:CH, (CH2)2, o-C6H4] in MeCN gave the diazabicyclooctanes II (X as before) in 46-77% yield. Irradiation of I [n=2, X=CH20] gave the

before) in 46-77% yield. Irradiation of I [n = 2, : azepine III in 46% yield. T 83095-08-3P 83095-09-4P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) RN 83095-08-3 HCAPLUS

SSUP-Pyrrolo[1,2-c]imidazol-5-one, 2-ethylhexahydro-7a-hydroxy-1-methyl-, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.

L69 ANSWER 47 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

83095-09-4 HCAPLUS 5H-Pyrrolo[1,2-c]imidazol-5-one, 2-ethylhexahydro-7a-hydroxy-1-methyl-, trans- [9CI] (CA INDEX NAME)

Relative stereochemistry.

L69 ANSWER 48 OF 63
ACCESSION NUMBER:
DOCUMENT NUMBER:
1982:7047 HCAPLUS
96:7047
Solvent effect in the reaction of (S)-N-isopropyl-Na-(benzyloxycarbonyl) prolinamide with lithium aluminum hydride
AUTHOR(S):
CORPORATE SOURCE:
SOURCE:
COMPORATE SOURCE:
COENT: CHITAG; ISSN: 0366-7022
DOCUMENT TYPE:
LANGUAGE:
GI

DOCUMENT TYPE: LANGUAGE: GI

- CONHCHMe2 - CR2NHCHMe2

AB A specific solvent effect in the reduction of the title amide (I) with LiALH4

H4

Was studied. The reactions in Et20 gave mainly II (R2 = 0) and II (R = H), while III and IV were produced in THF.

80090-65-0p

RL: SFN (Synthetic preparation); PREP (Preparation)
(preparation of)

80090-66-0 HCRPLUS

H-Pyrrol(1,2-c)imidazole, hexahydro-2-(1-methylethyl)-, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry

L69 ANSWER 49 OF 63
ACCESSION NUMBER:
DOCUMENT NUMBER:
SITTILE:
INVENTOR(S):
SOURCE:
SOURCE:
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT ANDOMETION:
FAMILY ACC. NUM. COUNT:
PATENT ANDOMETION:
TO SUBSTRANCE (S):
FAMILY ACC. NUM. COUNT:
PATENT ANDOMETION:
TO SUBSTRANCE (S):
FAMILY ACC. NUM. COUNT:
TO SUBSTRANCE (S)
FAMILY ACC. NUM. COUNT:
TO SUBSTRANCE (S)

LOS SUBSTRANCE (S)
FAMILY ACC. NUM. COUNT:
TO SUBSTRANCE (S)

LOS PATENT NUMBURION:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
DE 2728870	A1	19780112	DE 1977-2728870		19770627
DE 2728870	C2	19850411			
JP 53007620	A2	19780124	JP 1976-82321		19760710
JP 56005389	В4	19810204			
JP 53084915	A2	19780726	JP 1976-158645		19761230
JP 55046390	B4	19801122			
GB 1523090	A	19780831	GB 1977-24800		19770614
US 4150240	A	19790417	US 1977-813989		19770708
СН 630070	A	19820528	CH 1977-8493		19770708
PRIORITY APPLN. INFO.:			JP 1976-82321	A	19760710
			JP 1976-158645	A	19761230

GI

D-Penicillamine (I) was prepared by cleavage of II (R = H, PhCH2CO, PhOCH2CO: R1 = H, CO2H, CONH2, CONHPh, etc.) by aromatic amines. Thus, benzylpenicilloic acid  $\alpha$ -phenethylamide was heated with (PhNHCH2)2 in aqueous AcOH, followed by acidification with HCl to give 82.8% I.HCl. 66317-042. IT

66317-04-2
RL: RCT (Reactant); RACT (Reactant or reagent)
(ring cleavage of, by aromatic amines)
66317-04-2 HCAPLUS
Imidazo[5,1-b][thiazole-3,7-dicarboxylic acid, hexahydro-2,2-dimethyl-6(phenylmethyl)- (9CI) (CA INDEX NAME)

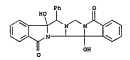


L69 ANSWER 50 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER: 1975:443238 HCAPLUS
DOCUMENT NUMBER: 83:43238
HITTLE: Photoreactions of bis(phthalimidomethyl)alkylamines
ROTH, H. J.; Schwarz, D.
CORPORATE SOURCE: Pharm. Inst., Univ. Bonn, Bonn, Fed. Rep. Ger.
SOURCE: Archiv der Pharmarie (Weinheim, Germany) (1975),
308(3), 218-24
CODEN: ARPPRAS; ISSN: 0365-6233
DOCUMENT TYPE: Journal
LANGUAGE: German
GI For diagram(s), see printed CA Issue.
AB The condensed imidazoles I and II (R = H, CHMe2, Ph) were obtained by photolysis of the amines III. Photolysis of III (R = Ph) also yielded
N-(3-hydroxyphthalimidinomethyl)phthalimide.
IT 56097-22-48 S0097-23-19 S0097-23-19
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
RN 56097-22-4 HCAPLUS
CN 6H, 13H, 15H-15oindolo[2",1":3',4"]imidazo[5',1":2,3]imidazo[5,1alisoindole-6,15-dione, 4b,4c,10b,11-tetrahydro-4b,10b-dihydroxy- (9CI)
(CA INDEX NAME)

RN 56097-25-7 HCAPLUS
CN 6H,13H,15H-Isoindolo[2'',1'':3',4']imidazo[5',1':2,3]imidazo[5,1a]isoindole-6,15-dione, 4b,4c,10b,11-tetrahydro-4b,10b-dihydroxy-l1-(1methylethyl)- (9CI) (CA INDEX NAME)

RN 56097-29-1 HCAPLUS 6H, 13H, 15H-1soindolo[2'',1'':3'',4']imidazo[5',1':2,3]imidazo[5,1-a]isoindol-e-6,15'-dione, 4b,4c,10b,11-tetrahydro-4b,10b-dihydroxy-11-phenyl-(9C1) (CA INDEX NAME)

L69 ANSWER 50 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Contin





L69 ANSWER 51 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1974:520538 HCAPLUS
DOCUMENT NUMBER: 81:120538
TITLE: Syntheses of imidazo[1,5-a] - and pyrazino[1,2-b]benzimidazoles
AUTHOR(S): Schubert, H.; Lettau, H.; Fischer, J.
CORPORATE SOURCE: Sekt. Chem., Martin Luther Univ., Halle, Ger. Dem.
Rep.
SOURCE: Tetrahedron (1974), 30(10), 1231-6
CODEN: TSTRAB; ISSN: 0040-4020

DOCUMENT TYPE: Journal
CODEN: TSTRAB; ISSN: 0040-4020

DOCUMENT TYPE: Journal
ANDIGAGE: German
OTHER SOURCE(S): CASREACT 81:120538
GI For diagram(s), see printed CA Issue.
AB 2-(a-aminobenzyl) benzimidazoles I (RI = H, cyclohexyl, CHZPh, alkyl, azyl). The benzhydryl analogs (II) were prepared similarly.
1,2-Dihydro-3H-imidazoles I (I), 5-a) benzimidazoles (III),
imidazo [1,5-a) benzimidazoles, 3-oxo-1,2,3,4-tetrahydropyrazino [1,2-a) benzimidazoles (IV), and 3,4-dioxo-1,2,3,4-tetrahydropyrazino [1,2-a) benzimidazoles were prepared by reaction of I and II with CH2O, COCI2, CLCHR2COCI, and (COC)12, resp. II (RI = H) with HC(OEt)3 gave
3,3-diphenyl-3H-imidazo[1,5-a) benzimidazole.

IT 3463-11-59 4463-12-69 5463-11-79
3463-11-59 54463-12-69 5463-11-79
3463-11-59 54463-12-69 5463-11-79
3463-11-59 FA463-11-59 54463-11-79
3463-11-5 HCAPLUS
CN 1H-Imidazo[1,5-a]benzimidazole, 2-cyclohexyl-2,3-dihydro-3-phenyl- (9CI) (CA INDEX NAME)

RN 54463-12-6 HCAPLUS
CN IN-Imidazo[1,5-a]benzimidazole, 2,3-dihydro-3-phenyl-2-(phenylmethyl)(SCI) (CA INDEX NAME)



RN 54463-13-7 HCAPLUS
CN IH-Imidazo[1,5-a]benzimidazole, 2,3-dihydro-2-(4-methoxyphenyl)-3-phenyl(9C1) (CA INDEX NAME)

54463-14-8 HCAPLUS 1H-Imidazo[1,5-a]benzimidazole, 2,3-dihydro-2-(4-methylphenyl)-3-phenyl-[GCI] (CA INDEX NAME)

54463-15-9 HCAPLUS 1H-Imidazo(1,5-a|benzimidazole, 2-(4-chlorophenyl)-2,3-dihydro-3,3-diphenyl-(9CI) (CA INDEX NAME)

54463-17-1 HCAPLUS 1H-Imidazo[1,5-a]benzimidazole, 2,3-dihydro-2-(4-nitrobenzoy1)-3,3-diphenyl- (9CI) (CA INDEX NAME)

54463-18-2 RCAPLUS 1H-Imidazo[1,5-a]benzimidazole, 2-cyclohexyl-2,3-dihydro-3,3-diphenyl-[9CI) (CA INDEX NAME)

L69 ANSWER 51 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

L69 ANSWER 52 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1974:437548 HCAPLUS
DOCUMENT NUMBER: 81:37548
Ethyl 1-oxoperhydropyrrolo {1,2-c} imidazolecarboxylates
INVENTOR(S): Fontanella, luigi: Occelli, Emilio
Gruppo Lepetit S.p.A.
SOURCE: Ger. Offen., 12 pp.
COODEN: GWXXEX
DOCUMENT TYPE: Patent
LANGUAGE: German
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PAIGNI NO.	KIND	DAIL	AFFBICATION NO.	DALL
DE 2354071	A1	19740516	DE 1973-2354071	19731029
GB 1381474	Α	19750122	GB 1973-45809	19731001
US 3901911	Α	19750826	US 1973-409985	19731026
FR 2205320	A1	19740531	FR 1973-38682	19731030
JP 49076893	A2	19740724	JP 1973-123938	19731102
PRIORITY APPLN. INFO.:			IT 1972-31275 A	19721103

GI For diagram(s), see printed CA 18000.

AB Ten esters [I; R = Me, Ph, C6H4OMe-4, or CH2Ph; Rl = rr, ...,

(OMe)
2-3,4], useful as anxiolytics, hypnotics, muscle relaxants, or sedatives,
were prepared by reaction of the pyrrolidines II with RICHO in the

were prepared by reaction of the pyrrolidines II with RICHO in the presence of 4-McG6H4SO3H in xylene at reflux, optionally followed by chromatog. separation into isomers.

IT 52840-79-6P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
RN 52840-79-6 HCAPLUS
CN H-Pyrrolo[1,2-c]imidazole-5-carboxylic acid, hexahydro-1-oxo-2-phenyl-, ethyl ester (9CI) (CA INDEX NAME)

L69 ANSWER 53 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1974:425642 HCAPLUS
DOCUMENT NUMBER: 81:25642 Synthesis of aryl-substituted 1,3- and 1,4-diazocine derivatives
AUTHOR(S): Sarges, Reinhard; Tretter, James R.
CORPORATE SOURCE: Cent. Res., Pfizer Inc., Groton, CT, USA
Journal of Organic Chemistry (1974), 39(12), 1710-16
CODEN: JOCEAH: ISSN: 0022-3263
JOURNAL SOURCE(S): CASREACT 81:25642
G1 For diagram(s), see printed CA Issue.
AB The synthesis of aryl-substituted 1,3- and 1,4-diazocine derivs. Was undertaken because their structural features suggested potential central nervous system activity. Reaction of Me B-(bromomethyl)cinnamate with N,M'-dimethylethylen-diamine gave Me N, N'-dimethyl-2phenylpiperazine-2-acetate which was converted to 1,4-dimethyl-7-phenyl1,2,3,4-tetrahydro-1,4-diazocin-5(BN)-one (I). Catalytic and hydride reduction of I led ultimately to the 6-phenylperhydro-1,4-diazocine (II). Conversion of trans-3-phenylproline to III followed by desulfurization and quaternization with MeI gave the bicyclic intermediate IV, which on

quaternization with MeI gave the bicyclic intermediate IV, which on treatment with NaH or Li-NH3 underwent transannular ring opening to give 1,3-dimethyl-6-phenyl-1,2,3,7-tetrahydro-1,3-diazocin-4[8H]-one (V) and its perhydro analog, resp. Reaction of IV with NaOMe or with NaBH4 led

peripheral ring cleavage giving N-methyl-3-phenylproline methyl ester and the corresponding alc., resp. 5:1212-44-59 5:1212-45-94 5:1212-46-59 RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 5:1212-44-3 HCAPUUS |
H-Pyrrolofi, 2-c]imidazol-1-one, hexahydro-2-methyl-7-phenyl-, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.

HCAPLUS HH-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-methyl-7-phenyl-, monohydrochloride, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

● HC1

51212-46-5 HCAPLUS
1H-Pyrrolo[1,2-c]imidazolium, hexahydro-2,4-dimethyl-1-oxo-7-phenyl-, iodide, trans- (9CI) (CA INDEX NAME)

L69 ANSWER 54 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

50613-39-3 HCAPLUS
L-threo-a-D-galacto-Octopyranoside, methyl 7-bromo-6,7,8-trideoxy-6-(tetrahydro-1-oxo-6-propyl-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)-1-thio-,[6(6R,7a3)]- (9CI) (CA INDEX NAME)

50613-40-6 HCAPLUS
D-erythro- $\alpha$ -D-galacto-Octopyranoside, methyl 6,8-dideoxy-6(tetrahydro-l-oxo-6-propyl-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)-1-thio-,
[6(6R,7as)]- (9CI) (CA INDEX NAME)

50613-43-9 HCAPLUS

Tethreo-a-D-galacto-Octopyranoside, methyl 7-chloro-6,7,8-trideoxy-6-(tetrahydro-1-oxo-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)-1-thio-, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L69 ANSWER 34 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1973:515852 HCAPLUS 79:115852 TITLE: Lincomycin-type compounds Argouelis, Alexander D.: Mager

79:115832 Lincomycin-type compounds Argoudelis, Alexander D.: Magerlein, Barney J. Upjohn Co. U.S., 17 pp. CODEN: USXXAM Patent

PATENT ASSIGNEE (S): SOURCE: DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

KIND A B4 PATENT NO. APPLICATION NO. DATE DATE 19710312 US 1971-123891 JP 1972-24440 US 1971-123891 US 3758454 JP 56047193 19730911 PRIORITY APPLN. INFO.: A 19710312

For diagram(s), see printed CA Issue.
Treatment of 1'-demethyllincomycin derivs, with RCHO (R = e.g. H, 2-furyl
p-BrCEH4) gave I (R = as above; Rl, R2, = H, alkyl, R3 = halo, OH, ONe),
useful as bactericides. Thus, 7(S)-chloro-7-deoxy-1'-demethyllincomycinHCl reacted with HCHO in aqueous NaOH to give I (R = H, R1 = Me, R2 =
R3 =

IT

50613-38-2 HCAPLUS

Lethreo-a-D-galacto-Octopyranoside, methyl 7-chloro-6,7,8-trideoxy-6-(tetrahydro-1-oxo-6-pentyl-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)-1-thio-, [6(6R,7a5]]- [9CI] (CA INDEX NAME)

L69 ANSWER 54 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

L69 ANSWER 55 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1973:147966 HCAPLUS
TITLE:
Chalcone derivatives
Okamoto, Kojin: Ishida, Ryuichi; Shintomi, Keiichi
PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
1
1
1
24TENT INCORMATION:
1
1
24TENT INCORMATION:
1
27366 HCAPLUS
COBLES
CALLUS

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. PATENT NO. KIND DATE DATE JP 48019595 B4 19730312 JP 1971-55101 19710722 JP 49032871 For diagram(s), see printed CA Issue. Thetitle compds. (I), antispasmodics and tranquilizers, were prepared by

AB Theticle compds. (I), antispasmodics and tranquilizers, were prepared the reaction of 1-oxopyrrolidinoimidazolidinyl-acylophenones with 1-oxopyrrolidinoimidazolidinylbenzaldehydes. E.g., 9.8 g p-(1-oxopyrrolidinoi[1,2-c]imidazolidin-2-yl)propiophenone an 9.3 g p-(1-oxopyrrolidinoi[1,2-c]imidazolidin-2-yl)propiophenone an 9.3 g p-(1-oxopyrrolidinoi[1,2-c]imidazolidin-2-yl)propiophenone an 9.3 g p-(1-oxopyrrolidinoi[1,2-c]imidazolidin-2-yl)propiophenone an 9.3 g heated 5 hr at 40-50° with 201 NaOH solution to give 721 [ (R1 = R3 = H, R2 = N6). Similarly prepared were 1.2HCl (R1, R2, R3 and % yield given):

Me, H, H, 58.5%; H, H, H, -: H, h, Me, 77.

1 41038-71-59 41124-24-79 41124-25-69

RL: SPN (Synthetic preparation): PREP (Preparation) (preparation of)

RN 41038-71-5 HCAPLUS
CN 1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-(4-[1-0xo-3-(4-(tetrahydro-1-oxo-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)phenyl]-2-propenyl]phenyl]-, dihydrochloride (9CI) (CA INDEX NAME)

### ●2 HC1

41124-24-7 HCAPLUS
1H-Pyrcolo[1,2-c]imidazol-1-one, 2,2'-[(2-methyl-3-oxo-1-propene-1,3-diyl)di-4,1-phenylene]bis[hexahydro-(9CI) (CA INDEX NAME)

ANSWER 55 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

41373-86-8 HCAPLUS
Benzaldehyde, 4-(tetrahydro-1-oxo-1H-pyrrolo(1,2-c)imidazo1-2(3H)-yl)-(SCI) (CA INDEX NAME)

32901-73-8 41373-89-1 41518-30-3
RL: RCT (Reactant): RACT (Reactant or reagent)
(reaction of, with benzaldehydes)
32901-73-8 HCRPLUS
H-Pyrrolo(1, 2-c)imidazol-1-one, 2-(4-acetylphenyl)hexahydro- (9CI) (CA INDEX NAME)

41373-89-1 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, 2-(4-acetyl-2-methylphenyl)hexahydro-(9C1) (CA INDEX NAME)

41518-30-3 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-[4-(1-oxopropyl)phenyl](9CI) (CA INDEX NAME)

L69 ANSWER 55 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 41124-25-8 HCAPLUS CN 1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-[4-[3-[3-methyl-4-(tetrahydro-

1-oxo-1H-pyrrolo[1,2-c]imidazol-2(3H)-yl)phenyl]-3-oxo-1-propenyl]phenyl], dihydrochloride (9CI) (CA INDEX NAME)

#### ●2 HC1

41373-84-6 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, 2,2'-[(1-methyl-3-oxo-1-propene-1,3-dyy)]di-4,1-phenylene]bis[hexahydro-, dihydrochloride (9CI) (CA INDEX NAME)

### ●2 HCl

32901-73-8 41373-86-8

32901-73-8 41373-86-8
RL: RCT (Reactant) rACT (Reactant or reagent)
(reaction of, with acetophenones)
32901-73-8 HCAPEUS
1H-Pytrolo[1,2-c]imidazol-1-one, 2-(4-acetylphenyl)hexahydro- (9CI) (CA
INDEX NAME)

L69 ANSWER 55 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN



L69 ANSWER 56 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1973:147963 HCAPLUS
DOCUMENT NUMBER: 7871:147963 HCAPLUS
TITLE: Chalcone derivatives
INVENTOR(S): OShiro, Susumu; Nagura, Takeo; Sugihara, Yukio;
Okamoto, Koji; Ishida, Ryuichi; Shintomi, Keiichi
Tanabe Seiyaku Co., Ltd.
Jpn. Rokai Tokkyo Koho, J pp.
CODEN: UKOMAF
DOCUMENT TYPE: Patent
LANGIJAGE: 7ANILY ACC. NUM. COUNT: 1

ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE KIND APPLICATION NO.

JP 48019594 B4 19730312 JP 1971-55100 19710722
JP 48012870 19740726
For diagram(s), see printed CA Issue.
The title compds. (I), antispassociates and tranquilizers, were prepared by treating 4,4'-diprolylaminochalcones with HCRO or with
N,N'-cathonyldimidazole (carbonalating agent). E.g., 18.5 g 4,4'-bis
(L-prolylamino)-B-methylchalcone in MeOH was stirred 5 hr at
50' with 13.4 g 37% HCHO to give 97% I (X = CH2, R1 = Me, R2 = R3 =
H). Similarly prepared were the following I (X, R1, R2, R3, and % yield given): CH2, H, Me, H, 85; CH2, H, H, H, 72% (dihydrochloride); CH2, H,

### ●2 HC1

41124-23-6 HCAPLUS 1H-Pyrrolo[1,2-c]midazol-1-one, 2,2'-[(l-methyl-3-oxo-1-propene-1,3-diylidi-4,1-phenylene]bis[hexahydro-(9CI) (CA INDEX NAME)

L69 ANSWER 57 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1973:97999 HCAPLUS DOCUMENT NUMBER: 78:97999 TITLE: N.N'-alkylidene pentides Parti

78:97999
N.N'-alkylidene peptides. Peptide synthesis by products in the action of carbonyl compounds Cardinaux, F.; Brenner, M. Inst. Org. Chem., Univ. Basel, Basel, Switz. Helvetica Chimica Acta (1973), 56(1), 339-47 CODEN: HACKAY; ISSN: 0018-019X

AUTHOR (S) :

CORPORATE SOURCE: SOURCE:

CODEN: HCACAV; ISSN: 0018-019X
JOURNAL
LANGUAGE:

Germa
AB Hydrogenolysis of Z-Pro-Leu-Glu(OCMe3)-Phe-OCMe3, Z-Val-His-Pro-PHeOMe,
and Z-Val-Tyr-Val-His-Pro-PHeOMe (Z = PhcH2O2C) yields by-products that
were identified as 4-imidazolidinone derivs. They were formed by
cycloaddn. of a carbonyl compound, formed by oxidation of the solvent
under the

The reaction conditions, to the newly liberated N-terminal of the peptide and to the N of the adjacent amino acid residue. 4019-118-019

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) 40149-18-6 HcAPUS L-Phenylalanne, N-[N-[4-methyl-1-oxo-2-(tetrahydro-1-oxo-1H-pyrrolo[1,2-c]imidazol-2(3M)-yl)pentyl]-L-a-glutaminyl]-, bis(1,1-dimethylethyl) ester, stereoisomer [9CI) (CA INDEX NAME)

L69 ANSWER 56 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

41124-24-7 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, 2,2'-[(2-methyl-3-oxo-1-propene-1,3-diyl)di-4,1-phenylene]bis[hexahydro-(9CI) (CA INDEX NAME)

RN 41124-25-8 HCAPLUS
CN 1H-Pyrrolo(1,2-c)imidazol-1-one,
hexahydro-2-[4-[3-[3-methyl-4-(tetrahydro-

1-oxo-1H-pyrrolo(1,2-c)imidazol-2(3H)-yl)phenyl]-3-oxo-1-propenyl}phenyl}, dihydrochloride (9CI) (CA INDEX NAME)

●2 HC1

L69 ANSWER 58 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1972:429613 HCAPLUS

TITLE: 17:29613 HCAPLUS

TITLE: Hydroxymethylation of antibiotics. VII.

Hydroxymethylation of N-demethylclindamycin

ACGRORATE SOURCE: ACGRORATE SOURCE: SOURCE: SOURCE: SOURCE: 10:2061 ACGRORATE SOURCE: COEN: JANTAJ; ISSN: 0021-8820

JOURNAL TYPE: JAURNAL COENTAGE

TOCIMENT TYPE: JAURNAL COENTAGE

ACCORN: JANTAJ; ISSN: 0021-8820

JOURNAL COENTAGE

JAURNAL COENTAGE

ACCORN: JANTAJ; ISSN: 0021-8820

DOCUMENT TYPE: Journal

(Continued)

```
L69 ANSWER 59 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1971:488591 HCAPLUS
DOCUMENT NUMBER: 75:88591
TITLE: 5-0xa-3,10-diazabicyclo[5.2.1]decane and
5-0xa-3,10-diazabicyclo[5.2.1]decan-4-one
derivatives
      potentially active on the central nervous system

PHOR(S): Fontanella, L.; Occelli, E.

Lab. Ric., Gruppo Lepetit S.p.A., Milan, Italy

WRCE: Farmaco, Edizione Scientifica (1971), 26(8), 685-709

CODEN: FRESAX; ISSN: 0430-0920

JOURNAT

HOUAGE: Italian

HER SOURCE(S): CASREACT 75:88591

For diagram(s), see printed CA Issue.

3,10-Diethyl-5-oxa-3,10-diazabicyclo[5.2.1] decane (I) and II are

spared
AUTHOR (S):
CORPORATE SOURCE:
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
            from III. Thus, III (R = R1 = Et) is treated with H2CO to give I. III
           = R1 = Me) is treated with COCl2 and KOH to give II (R = Me). Similarly prepared are 9 other II (R = C3-4 alkyl, PhCH2, Ph, (CH2)2NMe2, aralkyl,
           substituted phenyl).
33252-12-9P 33252-13-0P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
33252-12-9 HCAPEUS
H-Pytrolo[1,2-c]imidazole-5-methanol, hexahydro-2-methyl- (8CI) (CA INDEX NAME)
ΙT
                                         сн2-он
            33252-13-0 HCAPLUS
1H-Pyrrolo[1,2-c]imidazole-5-methanol, hexahydro-2-methyl-,
dihydrochloride (8CI) (CA INDEX NAME)
                                         сн2-он
```

●2 HC1

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

L69 ANSWER 60 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1971:449087 HCAPLUS
DOCUMENT NUMBER: 75:49087 Pyrrolidino[1,2-c]imidazolidinone derivatives
Oshiro, Susumu: Nakura, Takeo; Okamoto, Takashi;
Okumura, Kentato
Tanabe Seiyaku Co., Ltd.
OCUMENT TYPE: CODEN: JANXAD
PATENT INFORMATION: 1
PATENT INFORMATION: 1 PATENT NO. KIND DATE APPLICATION NO. DATE

JP 46016990 B4 19710511 JP 19680319

For diagram(s), see printed CA Issue.
I, useful as anti-inflammatory, analgesic, and antispasmodic drugs, are manufactured by reaction of II with R2CHO. II (R2 = Ph) (11.5 g) in 50 ml MeOH
is stirred 3 hr with 7.4 g 37% HCHO to give 11.7 g I (R1 = Ph, R2 = H),  L69 ANSWER 60 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued) ● HC1 32901-47-6 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, 2-(m-chlorophenyl)hexahydro- (8CI) (CA
INDEX NAME) 32901-48-7 HCAPLUS 1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-(p-methoxyphenyl)- (8CI) INDEX NAME) 32901-49-8 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-(p-methoxyphenyl)-,
monohydrochloride (8CI) (CA INDEX NAME)

L69 ANSWER 59 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN

Page 267

1

L69 ANSWER 60 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

N N Me

RN 32901-56-7 HCAPLUS
CN 1H-Pyrrolo(1,2-c)imidazol-1-one, hexahydro-2-(6-methyl-2-pyridyl)-, monohydrochloride (8CI) (CA INDEX NAME)

N N Me

• HC1

RN 32901-57-8 HCAPLUS
CN 1H-Pyrrolo]1,2-c|imidazol-1-one, hexahydro-2-propyl- (8CI) (CA INDEX NAME)

Pr-r

RN 32901-58-9 HCAPLUS
CN 1H-Pytrolo[1,2-c]imidazol-1-one, hexahydro-2-propyl-, monohydrochloride
(8C1) (CA INDEX NAME)

O Pr-n

• HCl

RN 32901-59-0 HCAPLUS

L69 ANSWER 60 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
CN 1H-Pyrrolo[1,2-c]imidazol-1-one, 2-(o-chlorophenyl)hexahydro- (8CI) (CA INDEX NAME)

RN 32901-64-7 HCAPLUS CN 1H-Pyrrolo[1,2-c]imidazol-1-one, 2-(o-chlorophenyl)hexahydro-, monohydrochloride (8CI) (CA INDEX NAME)

, N

• HCl

RN 32901-65-8 HCAPLUS
CN 1H-Pyrcolo[1,2-c]imidazol-1-one, 2-(p-chlorophenyl)hexahydro- (8CI) (CA INDEX NAME)

RN 32901-66-9 HCAPLUS
CN 1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-(o-methoxyphenyl)- (8CI)
(CA thirty name)

RN 32901-67-0 HCAPLUS
CN IH-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-(o-methoxyphenyl)-, monohydrochloride (8CI) (CA INDEX NAME)

L69 ANSWER 60 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)
CN 1H-Pyrrolo[1,2-c]imidazol-l-one, 2-butylhexahydro- (8CI) (CA INDEX NAME)

Bu-

RN 32901-60-3 HCAPLUS
CN 1H-Pyrrolo[1,2-c]imidazol-1-one, 2-butylhexahydro-, monohydrochloride
(8C1) (CA INDEX NAME)

Bu-n

HC1

RN 32901-61-4 HCAPLUS CN 1H-Pyrrolo[1,2-c]imidazol-1-one, 2-benzylhexahydro- (8CI) (CA INDEX NAME)

CH2-Ph

RN 32901-62-5 HCAPLUS
CN HH-Pyrcolofl,2-cjimidazol-1-one, 2-benzylhexahydro-, monohydrochloride
(8C1) (CA INDEX NAME)

CH2-Ph

HC1

RN 32901-63-6 HCAPLUS

169 ANSWER 60 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

• HCl

RN 32901-68-1 HCAPLUS CN HH-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-(m-nitrophenyl)- (8CI) (CA INDEX NAME)

NO2

RN 32901-69-2 HCAPLUS
CN IH-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-(m-nitrophenyl)-,
monohydrochloride (8CI) (CA INDEX NAME)

NO<sub>2</sub>

• HC1

RN 32901-70-5 HCAPLUS
CN IH-Pyrrolo(1,2-c)imidazol-1-one, hexahydro-2-(p-nitrophenyl)- (8CI) (CA INDEX NAME)

NO.

0

RN 32901-71-6 HCAPLUS
CN Benzeneaulfonamide, p-(tetrahydro-1-oxo-1H-pyrrolo[1,2-c]imidazo1-2(3H)-y1)- (8C1) (CA INDEX NAME)

32901-72-7 HCAPLUS
1H-Pyrrolo(1, 2-c]imidazol-1-one, 2-(m-aminophenyl)hexahydro-,
monohydrochloride (8CI) (CA INDEX NAME)

HC1

32901-73-8 HCAPLUS 1H-Pyrrolo[1,2-c]imidazol-1-one, 2-(4-acetylphenyl)hexahydro- (9CI) (CA

32901-74-9 HCAPLUS
1H-Pyrrolo[1, 2-c]imidazol-1-one, 2-(p-acetylphenyl)hexahydro-,
monohydrochloride (8CI) (CA INDEX NAME)

L69 ANSWER 60 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

● HC1

32902-37-7 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-phenyl- (8CI) (CA INDEX

32902-38-8 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, hexahydro-2-phenyl-, monohydrochloride
(8C1) (CA INDEX NAME)

• HC1

33035-95-9 RCAPLUS
1H-Pyrcolo[1,2-cp]imidazol-l-one, hexahydro-2-(p-nitrophenyl)-,
monohydrochloride (BCI) (CA INDEX NAME)

L69 ANSWER 60 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

• HCl

34062-99-2 HCAPLUS
1H-Pyrrolo[1,2-c]imidazol-1-one, 2-(p-chlorophenyl)hexahydro-,
monohydrochloride (8CI) (CA INDEX NAME)

L69 ANSWER 61 OF 63 HCAPLUS COPYRIGHT 2006 ACS ON STN

ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:
Reaction of 3-propylindole with aldehydes.
Preparation of 2-(a-aminoalkyl)indoles
Wolinsky, Joseph; Sundeen, J. E.
CORPORATE SOURCE:
SOURCE:
CORPORATE SOURCE:
COPEN: TETRAB; ISSN: 0040-4020
DOCUMENT TYPE:
LANGUAGE:
COPEN: TETRAB; ISSN: 0040-4020
THER SOURCE(S):
CASREACT 74:76254
GI For diagram(s), see printed CA Issue.
AB 2,2-\*2-Benzylidenebisindoles (I) are produced by the condensation of 3-propylindole with aromatic aldehydes. Aromatic aldehydes react initially, but reversibly, at N. N-Substituted products can be trapped as

as acetate derivs. and are converted to I under the original reaction conditions. 3-Propylindole reacts with HCHO and piperidine under mild conditions to give 1-piperidinemethyl-3-propylindole.

2-Piperidinemethyl-3-propylindole is obtained when the reaction with HCHO in AcOH is carried out at 100° in the presence of excess piperidine. The condensation of 3-propylindole with HCHO and primary amines, such as PhCHZNHZ, involves initial attack at N followed by intramol. substitution at the 2-position to yield

2-benzyl-2, 3-dihydro-9-propyl-1H-imidazo-[1,5-a]indole (II) and 2,4-dibenzyl-11-propyl-2,3,4,5-tetrahydro-1,3,5-triazepino[1,7-a]indole (III) Hydrolysis of cyclohexyl-Indiazoindole affords 2-cyclohexylaminomethyl-3-propylindole.

IT 30713-07-6P, 1H-Imidazo[1,5-a]indole, 2-cyclohexyl-2,3-dihydro-9-propyl- 30745-26-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 30713-07-6 HCAPLWS

CA

1H-Imidazo(1,5-a)indole, 2-cyclohexyl-2,3-dihydro-9-propyl- (9CI) (CA

30745-26-7 HCAPLUS 1H-Imidazo(1,5-a)indole, 2-benzyl-2,3-dihydro-9-propyl- (8CI) (CA INDEX

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L69 ANSWER 62 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1968:467279 HCAPLUS
DOCUMENT NUMBER: 69:67279
TITLE: This roll didne. I. Formation and reactions of
3,3'-methylenedithiazolidine
AUTHOR(S): Takatori, Toshisuke: Kojima, Masaharu; Taguchi,
Tanezo
CORPORATE SOURCE: Kyushu Univ., Fukuoka, Japan
SOURCE: Yakugaku Zasshi (1968), 88(3), 360-5
CODEN: YKKZAJ; ISSN: 0031-6903
DOCHMENT TYPE: Journal Journal
LANGUAGE: Journal
LANGUAGE: Journal
AB 2-Aminoethanethiol (3 g.) in 12 ml. H20 is treated with 3.5 ml. 378
formalin to give 3.1 g. 3,3'-methylenedithiazolidine (11 n = 0), m.
48-9' (petroleum ether). Similarly is prepared d1-3,3'-
methylenebis[perhydrocyclohexa(d[thiazole] (11 (n = 4); trans isomer m.
97-8'; cis isomer m. 101-2'. trans-2-Aminocyclohexanethiol
(2 g.) in 10 ml. H20 is treated with 2 ml. AcOH to give 1.5 g. trans-II
(R1 = Me, R2 = H), m. 50-1*; HCl salt m. 173-4'. Similarly
prepared are the following trans-II (R1, R2, m.p., and m.p. HCl salt
given):

Me, Me, - (b8 89-90'), 53-4', Ph, H, 53-4',
193-5'; and (R1R2 =) (CK215, -, 35-6', 225-7'.
Warming 2 g. 2,2'dithiazolidine in 10 ml. H20 with 2 ml. 378 formalin
gives perhydroimidazo[1,5-b:4,3-b']dithiazole (III), m. 80-1'
(petroleum ether). I (n = 0) (5.0 g.) and 3.0 g. PhOH is stirred 24 hrs.
in 20 ml. Et20 at room temperature, evaporated, and the residue in 1:1
petroleum ether: Store the member and the residue in 1:1
19505-80-7 HCAPLUS (H1)
RN 19505-80-7 HCAPLUS
CN 5H-Imidazo(5,1-b:4,3-b')bisthiazole, hexahydro- (7CI, 8CI) (CA INDEX
NAME)

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N N s

L69 ANSWER 63 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN (Continued)

4

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L69 ANSWER 63 OF 63 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1963:27236 HCAPLUS
SOCUMENT NUMBER: 58:27236
ORIGINAL REFERENCE NO.: 58:4536b-e
A made development in the formation reaction of thiazolidines from 2-arminoalkanethiols and carbonyl compounds
AUTHOR(S): A med development in the formation reaction of thiazolidines from 2-arminoalkanethiols and carbonyl compounds
AUTHOR(S): Taguchi, Tanezo: Takatori, Toshisuke: Kojima, Masaharu
CORPORATE SOURCE: Kyushu Univ., Fukuoka
CORDEN: CPBTAL; ISSN: 0009-2363
JOULIAN
AUGUAGE: Unavailable
OTHER SOURCE(S): CASEACT 58:27236
AB HSCHZCHZNH2 (I) treated with HCHO gave bis(3-thiazolidinyl) methane (II), m. 46-9°, which set free HCHO and gave the HCl salt (III) of thiazolidine (IV) with HCl-EtoH. The structure of II was confirmed by its infrared (I.R.) spectrum, and by its formation from IV with HCHO. I with other carbonyl compds, gave the corresponding simple thiazolidines, which did not give analogs of II with HCHO. 2, 2'-Dithiazolidine with HCHO gave perhydroimidazo[1,5-9:4,3-b']dithiazole (IV), m. 80-1*, confirmed by its I.R. spectrum. Similar treatment of the HCl salts of cis- and trans-aminocyclohexanethiols (VI) with HCHO gave cis-a and trans-cyclohexanethiols (VI) with HCHO gave (11-12°, resp., whereas free VI (like free I) gave snalogs of II, cis- and trans-3,3'-methylenebis/cyclohexaldthiazolidine), m. 101-2', and 97-8', resp., which, like II, freed HCHO and gave VII with HCH-EtoH. VII with D-glucose geok (by analogy with the products from I with D-glucose (Bonner and Meyer, CA 55, 13412b)] 2-(D-gluco-1,2,3,4,5-penthydroxypentyl)(-)-cis-and (-)-trans-cyclohexa[dithiazolidine] (IX), HCl salts m. 228-30' ([al170-cis-6) and (c)-place-1,2'-cyclohexa[dithiazolidine] (IX), HCl salts m. 228-30' ([al170-cis-6) and 229-31' ([al170-cis-6) and (c)-place-1,2'-cyclohexa[dithiazolidine] (IX), HCl salts m. 228-30' ([al170-cis-6) and (c)-prans-1) ([al170-des-6) and (c)-place-1,0'-cyclohexa[dithiazolidine] (IX), HCl salts m. 228-30' ([al170-cis-6) and (c)-place-1,0'-cy
```

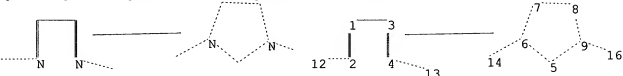
process claims

10/520,800

06/28/2006

=>

Uploading C:\Program Files\Stnexp\Queries\10520800\9.str



ring nodes:
5 6 7 8 9
ring/chain nodes:
1 2 3 4 12 13 14 16
ring/chain bonds:
1-2 1-3 2-12 3-4 4-13 6-14 9-16
ring bonds:
5-6 5-9 6-7 7-8 8-9
exact/norm bonds:

1-2 1-3 2-12 3-4 4-13 5-6 5-9 6-7 6-14 7-8 8-9 9-16

Connectivity:

2:2 E exact RC ring/chain 4:2 E exact RC ring/chain 5:2 E exact RC ring/chain Match level:
1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 12:CLASS 13:CLASS 14:CLASS 16:CLASS fragments assigned product role: containing 5 fragments assigned reactant/reagent role: containing 1 node mappings:
2:6 1:7 3:8 4:9 12:14 13:16

L60 STRUCTURE UPLOADED

=> s L60

SAMPLE SEARCH INITIATED 12:16:23 FILE 'CASREACT'
SCREENING COMPLETE - 7817 REACTIONS TO VERIFY FROM

611 DOCUMENTS

1 DOCS

64.0% DONE 5000 VERIFIED 2 HIT RXNS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*
BATCH \*\*COMPLETE\*\*
PROJECTED VERIFICATIONS: 151073 TO 161607
PROJECTED ANSWERS: 1 TO 141

L61 1 SEA SSS SAM L60 ( 2 REACTIONS)

=> s L60 full

FULL SEARCH INITIATED 12:16:30 FILE 'CASREACT'

SCREENING COMPLETE - 169853 REACTIONS TO VERIFY FROM 12466 DOCUMENTS

100.0% DONE 169853 VERIFIED 111 HIT RXNS ( 3 INCOMP) 35 DOCS

SEARCH TIME: 00.00.02

L62 35 SEA SSS FUL L60 ( 111 REACTIONS)

=> d sca

=> file casreact
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ENTRY SESSION
332.05 1388.13

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

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ENTRY SESSION
-47.25 -47.25

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FILE CONTENT: 1840 - 25 Jun 2006 VOL 144 ISS 26

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Some CASREACT records are derived from the ZIC/VINITI database (1974-1991) provided by InfoChem, INPI data prior to 1986, and Biotransformations database compiled under the direction of Professor Dr. Klaus Kieslich.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d his

(FILE 'HOME' ENTERED AT 09:32:44 ON 28 JUN 2006)

FILE 'REGISTRY' ENTERED AT 09:32:52 ON 28 JUN 2006 L1 1650308 S NCNC2/ESS

FILE 'REGISTRY' ENTERED AT 09:51:45 ON 28 JUN 2006
L2 STRUCTURE UPLOADED
L3 8980 S L2 FULL
L4 STRUCTURE UPLOADED
L5 3519 S L4 FULL

L6 3519 S L4 FOLL L6 50 S L4

L7 50 S L2 SAVE TEMP L5 GLOR800STR2/A

FILE 'HCAPLUS' ENTERED AT 10:08:22 ON 28 JUN 2006

L8 253 S L5

FILE 'REGISTRY' ENTERED AT 10:08:52 ON 28 JUN 2006 L9 628 S NC>1 AND L5

FILE 'STNGUIDE' ENTERED AT 10:12:24 ON 28 JUN 2006

FILE 'REGISTRY' ENTERED AT 10:14:37 ON 28 JUN 2006

- L10 STRUCTURE UPLOADED
- L11 50 S L10 SAM SSS SUB=L5
- L12 1426 S L10 FULL SSS SUB=L5
- FILE 'HCAPLUS' ENTERED AT 10:17:32 ON 28 JUN 2006 L13 108 S L12
- FILE 'REGISTRY' ENTERED AT 10:17:46 ON 28 JUN 2006 L14 237 S L12 AND NRRS>2
- FILE 'HCAPLUS' ENTERED AT 10:27:31 ON 28 JUN 2006 L15 56 S L14
- FILE 'REGISTRY' ENTERED AT 10:27:55 ON 28 JUN 2006 L16 1189 S L12 NOT L14
- FILE 'HCAPLUS' ENTERED AT 10:28:11 ON 28 JUN 2006
- L17 60 S L16
- L18 8 S L15 AND L17
  - FILE 'REGISTRY' ENTERED AT 10:29:05 ON 28 JUN 2006
  - FILE 'STNGUIDE' ENTERED AT 10:29:18 ON 28 JUN 2006
- FILE 'REGISTRY' ENTERED AT 10:45:47 ON 28 JUN 2006
- L19 STRUCTURE UPLOADED
- L20 32 S L19 SAM SSS SUB=L12
- L21 551 S L19 FULL SSS SUB=L12
- FILE 'HCAPLUS' ENTERED AT 10:52:12 ON 28 JUN 2006 L22 85 S L21
- FILE 'REGISTRY' ENTERED AT 10:53:23 ON 28 JUN 2006 L23 368 S L21 NOT L14
- FILE 'HCAPLUS' ENTERED AT 10:53:51 ON 28 JUN 2006 L24 45 S L23
- FILE 'REGISTRY' ENTERED AT 10:54:25 ON 28 JUN 2006
- L25 875 S L12 NOT L21
- L26 821 S L12 NOT (L21 OR L14)
- L27 3310 S 180.306.6/RID
- L28 809 S L26 AND L27
- L29 12 S L26 NOT L28
  - FILE 'STNGUIDE' ENTERED AT 11:02:16 ON 28 JUN 2006
- FILE 'REGISTRY' ENTERED AT 11:04:38 ON 28 JUN 2006
- L30 STRUCTURE UPLOADED
- L31 1 S L30 SAM SSS SUB=L12
- L32 35 S L30 FULL SSS SUB=L12
- L33 0 S L32 AND L24
- L34 35 S L32 AND L14

FILE 'HCAPLUS' ENTERED AT 11:07:03 ON 28 JUN 2006 L35 11 S L34 FILE 'REGISTRY' ENTERED AT 11:07:31 ON 28 JUN 2006 FILE 'HCAPLUS' ENTERED AT 11:08:14 ON 28 JUN 2006 L36 54 S L35 OR L24 FILE 'REGISTRY' ENTERED AT 11:11:52 ON 28 JUN 2006 L37 403 S L23 OR L32 FILE 'HCAPLUS' ENTERED AT 11:13:36 ON 28 JUN 2006 L38 1 S US2005-520800/APPS SEL RN FILE 'REGISTRY' ENTERED AT 11:14:26 ON 28 JUN 2006 L39 109 S E1-E109 L40 34 S L39 AND L37 L41 75 S L39 NOT L40 L42 9 S L14 AND L39 L43 202 S L14 NOT L32 FILE 'HCAPLUS' ENTERED AT 11:26:41 ON 28 JUN 2006 L44 47 S L43 FILE 'REGISTRY' ENTERED AT 11:27:00 ON 28 JUN 2006 FILE 'STNGUIDE' ENTERED AT 11:31:48 ON 28 JUN 2006 FILE 'REGISTRY' ENTERED AT 11:35:19 ON 28 JUN 2006 STRUCTURE UPLOADED L45 L46 2 S L45 SAM SSS SUB=L12 46 S L45 FULL SSS SUB=L12 L47 46 S L14 AND L47 L48 FILE 'HCAPLUS' ENTERED AT 11:37:27 ON 28 JUN 2006 L49 12 S L48 L50 63 S L24 OR L35 OR L49 FILE 'REGISTRY' ENTERED AT 11:38:49 ON 28 JUN 2006 L51 191 S L14 NOT L47 L52 161 S L14 NOT (L47 OR L32) FILE 'REGISTRY' ENTERED AT 11:52:20 ON 28 JUN 2006 SAVE TEMP L23 GLOR800L23/A SAVE TEMP L34 GLOR800L34/A SAVE TEMP L48 GLOR800L48/A FILE 'HCAPLUS' ENTERED AT 11:55:09 ON 28 JUN 2006 SAVE TEMP L50 GLOR800L50/A FILE 'CASREACT' ENTERED AT 12:03:57 ON 28 JUN 2006 L53 STRUCTURE UPLOADED

> 1 S L53 SAM SSS 113 S L53 FULL SSS

> > STRUCTURE UPLOADED

85 S L55/COM

L54

L55

L56 L57

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1 S L57 SAM SSS
L58
L59
           8 S L57 FULL SSS
            STRUCTURE UPLOADED
L60
L61
           1 S L60
L62
          35 S L60 FULL
   FILE 'HCAPLUS' ENTERED AT 12:20:00 ON 28 JUN 2006
L63
      35 S L62
L64
           3 S L50 AND L63
L65
           24 S GLORIUS F?/AU
           6 S L65 AND L50
L66
L67
           4 S L65 AND L63
L68
           3 S L66 AND L67
           63 S L66 OR L50
L69
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FILE 'REGISTRY' ENTERED AT 12:26:44 ON 28 JUN 2006

FILE 'HCAPLUS' ENTERED AT 12:26:50 ON 28 JUN 2006

FILE 'CASREACT' ENTERED AT 12:29:25 ON 28 JUN 2006

 $\Rightarrow$  d ibib abs hit L62 1-35

```
L62 ANSWER 1 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 144:350821 CASREACT
ITITLE: Influence of annelation in N-heterocyclic carbenes:
Novel quinoxaline-annelated NHCs trapped as
                                                                                                                                                                                                                                                       L62 ANSWER 1 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                                                                                                                                                                                                                                                                                                                                                (Continued)
                                                                    metal complexes
Saravanakumar, Shanmuganathan: Kindermann, Markus K.:
Heinicke, Joachim: Koeckerling, Martin
Inatitut fuer Chemie und Biochemie,
Ernst-Moritz-Arndt-Universitaet Greifawald,
Greifawald, 17487, Germany
Chemical Communications (Cambridge, United Kingdom)
(2006), (61), 640-642
CODEN: CHCOFS: ISSN: 1359-7345
Royal Society of Chemiatry
Journal
  transition
  AUTHOR (S):
  CORPORATE SOURCE:
  SOURCE:
  PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
                                                                                                                                                                                                                                                        G: CM 1
YIELD 52%
                                                                                                                                                                                                                                                                                                      G: CM 2
YIELD 52%
               UAGE: English
Quinoxaline-annelated imidazol-2-ylidenes are less stable in comparison
                                                                                                                                                                                                                                                                                  RCT C 881020-50-4, F 122-51-0
RGT H 16941-11-0 PF6.NH4
PRO G 881020-53-7
               their non-annelated analogs, featuring high acidity of the C-H group of the parent quinoxalino[2,3-d]imidazolium salts: rhodium and silver complexes of the quinoxalino[2,3-d]imidazolium salts: rhodium and silver complexes of the quinoxalino[2,3-d]imidazolylidenes were isolated and characterized. Orthoformate condensation with N.N'-R2-2,3-quinoxalinediamine gave 1,3-R2-quinoxalino[2,3-d]imidazolium hexafluorophosphates (2a,b; R = tBuCH2, iPr,) Deprotonation of 2a by KH in the presence of [Rh[cod]Cl]2 gave the corresponding ([h-CHZED]HRh[cod]Cl] [h-CHZED = 1,3-dlimopenty]quinoxalino[2,3-d]imidazol-2-ylidene], whereas the free ligand L-CHZED (3) is unstable and non-detectable even at -50'. Metalation of 2b y Ag2O gave the cationic silver complex [[L-iPr]ZAg]PF6. The synthesis, NMR-, and cal
                                                                                                                                                                                                                                                        RX (3)
                                                                                                                                                                                                                                                                                               122-51-0 CH(OEt)3
5 hours, 120 deg C
                                                                                                                                                                                                                                                       RX(4) OF 18
                                                                                                                                                                                                                                                                                                    ...E + F ===> I...
crystal
structure data of novel electron-deficient quinoxaline annelated imidazol-2-ylidene precursors and complexes thereof are reported and compared with related less electron-withdrawing or non-annelated N-heterocyclic carbenes and complexes to illustrate annelation effects.

REFERENCE COUNT: 38 THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS
                                                                                                                                                                                                                                                                                                                                                         Et
                                                                                                                                                                                                                                                                                                                                                                                 <del>(4)</del>>
                                                                                     RECORD. ALL CITATIONS AVAILABLE IN THE RE
 FORMAT
                                                                                                                                                                                                                                                                                                                                                                                                           I: CM 1
YIELD 76%
                                             ...C + F ===> G...
 RX(3) OF 18
                                                            CMea
                                                                                                                                                                                                                                                       I: CM 2
YIELD 76%
                                                                                                                                    (3)
 c
                                                                                F
L62 ANSWER 1 OF 35 CASREACT COPYRIGHT 2006 ACS on STN RX(4) RCT E 881020-51-5, F 122-51-0 RGH 16941-11-0 PF6.NH4 PRO I 881020-55-9 SOL 122-51-0 CH(OEL)3 CON 24 hours, 120 deg C
                                                                                                                                                                         (Continued)
                                                                                                                                                                                                                                                       L62 ANSWER 1 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                           Q + R + S ===> T...
 RX(7) OF 18
                                                                                                                                                       ● Aq(I)
                                                                                                                                                                                                                                                       V: CM 1
YIELD 71%
                                                                                                                                                                                                                                                                                                      V: CM 2
YIELD 71%
                                                                                                                                                                                                                                                                                  RCT
RGT
PRO
SOL
                                                                                                                                                                                                                                                                                             Q 78198-90-0, R 18997-19-8
W 14104-20-2 AgBF4
V 881020-59-3
75-09-2 CH2C12
                                                                                                                                                                                                                                                       RX (9)
                          T: CM 1
YIELD 72%
                                                                                                                                                                                                                                                                                                                          50 deg C
                                                                                                                                                                                                                                                       RX(15) OF 18 COMPOSED OF RX(7), RX(8) RX(15) Q + R + S ===> U
                                                                                                                                                                                                                                                       Me 3C
                                                                                                                                                                                                                                                                                                                                               C1
 T: CM 2
YIELD 72%
                                                                                                                                                                                                                                                                                                                                                                                                             ● Aq(I)
                            RCT Q 78198-90-0, R 18997-19-8, S 2923-28-6
PRO T 883990-73-6
SOL 75-09-2 CH2C12
CON 24 hours, 50 deg C
NTE in the dark
```

RX(9) OF 18

Q + R ===> V...

L62 ANSWER 1 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

U YIELD 55%

RCT Q 78198-90-0, R 18997-19-8, S 2923-28-6 PRO T 883990-73-6 SOL 75-09-2 CH2C12 CON 24 hours, 50 deg C NTE in the dark RX (7)

RX (8)

RCT RGT PRO

T 883990-73-6 L 7693-26-7 KH U 881020-60-6 109-99-9 THF overnight, -78 deg C -> room temperature

L62 ANSWER 2 OF 35
ACCESSION NUMBER:
11712:

CORPORATE SOURCE:

SOURCE:

COMPORATE SOURCE

PUBLISHER: Royal Society of Chemistry
DOCUMENT TYPE: Journal
LANGUAGE: English
AB The donor properties of aryl-substituted N-heterocyclic carbenes (NHC)
were characterized by lone pair donation from the carbene C and, as is
shown here, by donation of electron d. of the aromatic x-face of the NHC
aryl groups towards the metal. The variation of the remote substituents
R

arys groups towards the metal. The variation of the remote substituents

(R = H, OC12H25, Me, Br) on the Ph ring of ruthenium diphenyl-substituents
imidazolylidene-based NMC complexes has a significant influence on the
redox behavior of these Grubbs II and Grubbs-Hoveyda type metathesis
catalysts, and can be used to modify the catalytic activity of such
complexes. As evidenced by cyclic voltammetric studies of Grubbs-Hoveyda
type complexes, the saturated and unsatd. NMC ligands can give rise to
different redox potentials Ru(II)/Ru(III). The systematic changes of the
redox potential according to the electron-donating nature of the remote
substituents and the fact that the aryl ring is electronically decoupled
from the N heterocycles provides strong evidence of the x-face
coordination of the Ru-carbene.

REFERENCE COUNT: 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR
THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

RX (13) OF 102

...AC + AB ==> I...

L62 ANSWER 2 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

• c1

RCT AC 181709-91-1, AB 64-18-6 RGT AD 7647-01-0 HC1 PRO I 221154-71-9 SOL 123-91-1 Dioxane NTE conditions not stated RX (13)

RX (26) OF 102 ...AH + AB ===> A...

L62 ANSWER 2 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

AM 56222-36-7, AB 64-18-6 AD 7647-01-0 HCl A 160256-31-5 123-91-1 Dioxane conditions not stated RX (26)

RX(27) OF 102 ...AR + AB ===> AV...

(Continued)

L62 ANSWER 2 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

Me (CH2) 11 (CH<sub>2</sub>)11

ΑV

RCT AK 974184-80-2, AB 64-18-6 RGT AD 7647-01-0 HCl PRO AV 974184-82-4 SOL 123-91-1 Dioxane NTE conditions not stated RX (27)

RX(42) OF 102 COMPOSED OF RX(16), RX(29) RX(42) AH + AA ===> AW

AW

RCT AH 49673-43-0 RGT AJ 16853-85-3 LiAlH4 PRO AI 475578-15-5 SOL 109-99-9 THF NTE conditions not stated RX (16)

RCT AI 475578-15-5, AA 122-51-0 RGT AB 64-18-6 HCO2H PRO AW 411245-49-8 NTE solvent and conditions not stated RX (29)

RX(43) OF 102 COMPOSED OF RX(17), RX(30) RX(43) AR + AA ===> AX

L62 ANSWER 2 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

L62 ANSWER 2 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

AX

RCT AK 874184-80-2 RGT AJ 16853-85-3 LIA1H4 PRO AL 874184-78-8 SOL 109-99-9 THF NTE conditions not stated RX (17)

AL 874184-78-8, AA 122-51-0 AB 64-18-6 HCO2H AX 874184-84-6 solvent and conditions not stated RX (30)

RX(44) OF 102 COMPOSED OF RX(18), RX(31) RX(44) AN + AA ===> G

L62 ANSWER 2 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

RCT AM 56222-36-7 RGT AJ 16853-85-3 LIA1H4 PRO AN 258278-23-8 SOL 109-99-9 THF NTE conditions not stated RX (18) RCT AN 258278-23-8, AA 122-51-0 RGT AB 64-18-6 HCO2H PRO G 245679-17-8 WTE solvent and conditions not stated RX (31)

RX(45) OF 102 COMPOSED OF RX(19), RX(12) RX(45) AC + AA ===> K

L62 ANSWER 2 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

STEPS

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RCT AC 181709-91-1 RGT AJ 16853-85-3 LiAlH4 PRO 2 870123-16-3 SOL 109-99-9 THF NTE conditions not stated

Z 870123-16-3, AA 122-51-0 AB 64-18-6 HCO2H K 874184-69-7 solvent and conditions not stated

RX(46) OF 102 COMPOSED OF RX(20), RX(32) RX(46) AO + AA ===> AF

STEPS

L62 ANSWER 2 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

AГ

AO 874184-81-3 AJ 16853-85-3 LiAlH4 AP 874184-79-9 109-99-9 THF conditions not stated RX (20)

AP 874184-79-9, AA 122-51-0 AB 64-18-6 HCO2H AF 874184-85-7 solvent and conditions not stated RX (32)

L62 ANSWER 3 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 143:439653 CASREACT
TITLE: Room-Temperature Negishi Cross-Coupling of
Unactivated Alkyl Bromides with Alkyl Organozinc Reagents Utilizing a Pd/N-Heterocyclic Carbene Catalyst Hadel, Niloufar; Kantchev, Eric Assen B.; O'Brien, Christopher J.; Organ, Michael G. Department of Chemistry, York University, Toronto, AUTHOR (S): CORPORATE SOURCE: M3J 1P3, Can. Journal of Organic Chemistry (2005), 70(21), SOURCE: 8503-8507 CODEN: JOCEAH: ISSN: 0022-3263 American Chemical Society Journal English PUBLISHER: DOCUMENT TYPE: LANGUAGE: AB A high-yie UAGE: English
A high-yielding cross-coupling reaction of unactivated alkyl bromides
possessing B-hydrogens with alkylzinc halides utilizing a
Pd/N-heterocyclic carbene (NHC) catalyst at room temperature is PG/N-neterocyclic carpene (NHC) Gatalyst at from temperature is variety of Pd sources, Pd2(dba)3, Pd(OAc)2, or PdBr2, with the comavailable ligand precursor 1,3-bis(2,6-diisopropylphenyl)imidazolium chloride successfully coupled 1-bromo-3-phenylpropane with n-butylzinc bromide in THF/NMP. An investigation of different NHC precursors showed that the bulky 2,6-diisopropylphenyl moiety was necessary to achieve high coupling yields (75-851). The corresponding Et analog was moderately active (11%). A range of unsym. NHC precursors were prepared and evaluated. active (11%). A range of unsym. NHC precursors were prepared and evaluated.

The ligand precursor containing one 2,6-disopropylphenyl and one 2,6-diethylphenyl afforded the coupling product in 47% yield, clearly suggesting a direct relationship between the steric topog. created by the flanking N-substituents and caralyst activity. Under optimal conditions, a number of alkyl bromides and alkylzinc halides possessing common functional groups (amide, nitrile, ester, acetal, and alkyne) were effectively coupled (61-92%). It is noteworthy that B-substituted alkyl bromides and alkylinch halides successfully underwent cross-coupling. Also, under these conditions alkyl chlorides were unaffected.

REFERENCE COUNT: 70 THERE ARE 70 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Eto

...C + F ===> G

• c1 G YIELD 56% RX (2) RCT C 118923-23-2 STAGE(1) RGT H 14044-65-6 BH3-THF SOL 109-99-9 THF CON 18 hours, reflux STAGE(2)

RGT I 67-56-1 MeOH

CON room temperature STAGE(3) RCT F 122-51-0 RCT J 7647-01-0 HC1 SOL 7732-18-5 Water, 122-51-0 CH(OEt)3 CON 2 hours, 120 deg C PRO G 868593-18-4 RX (8) OF 34 ...S + F ==> AA

Eto

L62 ANSWER 3 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

(8)

(Continued)

L62 ANSWER 3 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

Et Et H i-Pr

• (1)

AA YIELD 78%

RX(8) RCT S 868593-23-1

STAGE(1)

RGT H 14044-65-6 BH3-THF

SOL 109-99-9 THF

CON 18 hours, reflux

STAGE(2) RGT I 67-56-1 MeOH CON room temperature

STAGE(3)
RCT F 122-51-0
RGT J 7647-01-0 HC1
SOL 7732-18-5 Water, 122-51-0 CH(OEt)3
CON 2 hours, 120 deg C

PRO AA 866926-58-1

RX(9) OF 34 ...V + F ===> AB

L62 ANSWER 3 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued PRO AB 866926-59-2

• (

AC YIELD 82%

RX(10) RCT X 868593-27-5

STAGE(1)

ROT H 14044-65-6 BH3-THF

SOL 109-99-9 THF

CON 24 hours, reflux

STAGE(2)

ROT I 67-56-1 MeOH

CON room temperature

STAGE(3)

ROT F 122-51-0

ROT J 7647-01-0 HC1

SOL 7732-18-5 Water, 122-51-0 CH(OEt)3

i-Pr Pr-i

H P Me Me Me

● C1 -AB YIELD 52%

RX(9) RCT V 868593-25-3

STAGE(1)

RGT H 14044-65-6 BH3-THF

SOL 109-99-9 THF

CON 18 hours, reflux

STAGE(2)

RGT I 67-56-1 MeOH

CON room temperature

STAGE(3)

RCT F 122-51-0

RGT J 7647-01-0 HC1

SOL 7732-18-5 Water, 122-51-0 CH(OEt)3

CON 2 hours, 120 deg C

L62 ANSWER 3 OF 35 CASREACT COPYRIGHT 2006 ACS on STN CON 2 hours, 120 deg C

PRO AC 868593-33-3

RX(11) OF 34 ...E + F ===> AD

F--5+ F--F-AD: CM 1 YIELD 378

AD: CM 2 YIELD 378

RX(11) RCT Z 868593-29-7

STAGE(1)
RGT H 14044-65-6 BH3-THF
SOL 109-99-9 THF
CON 24 hours, reflux

STAGE(2)
RGT I 67-56-1 MeOH
CON room temperature

STAGE(3)
RCT F 122-51-0
RGT AE 16941-11-0 PF6.NH4
SOL 109-99-9 THF
CON 18 hours, 80 deg C

(Continued)

L62 ANSWER 3 OF 35 CASREACT COPYRIGHT 2006 ACS on STN PRO AD 868593-36-6 (Continued) L62 ANSWER 4 OF 35
ACCESSION NUMBER:
143:172810 CASREACT
Inidazo[1,3-a]pyridine-3-ylidenes-pyridine derived
N-heterocyclic cachene ligands
Frank
CORPORATE SOURCE:
MAX-Planck-Institut fuer Kohlenforschung, Muelheim an
der Ruhr, 45470, Germany
Tetrahedron (2005), 61(25), 6207-6217
CODEN: TETRAB: ISSN: 0040-4020
Lisevier B.V.
Journal
LANGUAGE:
English

AB The ready synthesis of differently substituted
2H-imidazo[1,5-a]pyridin-4
ium bromides, e.g., I, is reported. These salts were precursors for a
class of N-heterocyclic carbene ligands. As a consequence of their
bicyclic geometry, these ligands are capable of influencing the
coordination sphere of a carbene bound metal. The usefulness of these
ligands was demonstrated in the palladium-catalyzed Suzuki-Miyaura
cross-coupling of sterically hindered aryl chlorides.

REFERENCE COUNT: 31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR
THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

...Y + Z + Q ===> AA RX (9) OF 53

L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

RX (9) RCT Y 18997-19-8, Z 2923-28-6 STAGE(1)
SOL 75-09-2 CH2Cl2
CON 45 minutes, room temperature STAGE(2)
RCT Q 861404-00-4
CON SUBSTAGE(1) 19 hours, 40 deg C
SUBSTAGE(2) 40 deg C -> room temperature STAGE (3) SOL 67-56-1 MeOH STAGE (4)

RGT AB 1643-19-2 Bu4N.Br

SOL 75-09-2 CH2Cl2

CON 2 hours, room temperature PRO AA 861404-15-1 NTE in the dark

RX(10) OF 53 ...Y + Z + T ===> AD

L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (10)

AD: CM 2 YIELD 52%

RX (10) RCT Y 18997-19-8, Z 2923-28-6

STAGE(1)
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature STAGE(2)
RCT T 861404-01-5
CON SUBSTAGE(1) 24 hours, 40 deg C
SUBSTAGE(2) 40 deg C -> room temperature STAGE (3) SOL 64-17-5 EtOH STAGE(4)

RGT AB 1643-19-2 Bu4N.Br

SOL 75-09-2 CH2C12

CON 2 hours, room temperature PRO AD 861404-16-2 NTE in the dark

RX(11) OF 53 ...Y + Z + V ===> AE

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L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN MTE in the dark
L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                              (Continued)
                                                                                                                                                                                                                                         (Continued)
                                                                                                                                                                 ...Y + W ===> AF
                                                                                                                                          RX (12) OF 53
                                                                                                                                                                                                                       (12)
(11)
AE: CM 1
YIELD 47%
RX(11) RCT Y 18997-19-8, Z 2923-28-6
                                                                                                                                          AF
YIELD 22%
                 STAGE(1)
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature
                                                                                                                                                         RCT Y 18997-19-8
                                                                                                                                          RX (12)
                                                                                                                                                            STAGE(1)
RGT Z 2923-28-6 Ag03SCF3
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature
                 STAGE(2)
RCT V 861404-02-6
CON SUBSTAGE(1) 14 hours, 40 deg C
SUBSTAGE(2) 40 deg C -> room temperature
                                                                                                                                                            STAGE(2)
RCT W 851404-03-7
CON SUBSTAGE(1) 20 hours, 45 deg C
SUBSTAGE(2) 45 deg C -> room temperature
                 STAGE(3)
SOL 64-17-5 EtOH
                 STAGE (4)

RGT AB 1643-19-2 Bu4N.Br

SOL 75-09-2 CH2C12

CON 12 hours, room temperature
                                                                                                                                                            STAGE (3)
SOL 64-17-5 EtOH
CON room temperature
               PRO AE 861404-18-4
L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN PRO AF 861404-08-2 NTE in the dark
                                                                                              (Continued)
                                                                                                                                          L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                                                                                                                                                         (Continued)
                                                                                                                                           RX(33) OF 53 COMPOSED OF RX(13), RX(14)
RX(33) Y + X + AH ===> AI
RX(13) OF 53
                      ...Y + X ===> AG...
                                                                                                                                                                                                                       АН
                                                                            (13)
                                                                                                                                          STEPS
AG
YIELD 54%
                                                                                                                                          AI
YIELD 97%
              RCT Y 18997-19-8
RX (13)
                 STAGE(1)
RGT Z 2923-28-6 AgO3SCF3
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature
                                                                                                                                          RX (13)
                                                                                                                                                          RCT Y 18997-19-8
                                                                                                                                                             STAGE(1)

RGT 2 2923-28-6 AgO3SCF3

SOL 75-09-2 CH2C12

CON 45 minutes, room temperature
                 STAGE(2)
RCT X 861404-04-8
CON SUBSTAGE(1) 17 hours, 45 deg C
SUBSTAGE(2) 45 deg C -> room temperature
                                                                                                                                                             STAGE (2)
RCT )
CON 5
                                                                                                                                                                        )
X 861404-04-8
SUBSTAGE(1) 17 hours, 45 deg C
SUBSTAGE(2) 45 deg C -> room temperature
                 STAGE(3)
SOL 64-17-5 EtOH
CON room temperature
                                                                                                                                                             STAGE(3)
SOL 64-17-5 EtOH
CON room temperature
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PRO AG 661404-09-3 NTE in the dark

(Continued)

L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

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L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)
PRO AG $61404-09-3
NTE in the dark

RX(14) RCT AG $61404-09-3

STAGE(1)
CAT 14221-01-3 Pd(PPh3)4
SOL 110-71-4 (CH2OMe)2
CON 30 minutes, room temperature

STAGE(2)
RCT AM 4363-35-3
RGT AM 497-19-8 Na2CO3
SOL 7732-18-5 Water
CON 25 hours

STAGE(3)
SOL 7732-18-5 Water
CON room temperature

PRO AI $61404-11-7

RX(34) OF 53 COMPOSED OF RX(13), RX(15)
RX(34) Y + X + AN ===> AN

Br
He
He
He
```

\*\*STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*\*

RX(13) RCT Y 18997-19-8

\*\*STAGE(1)

RGT Z 2923-28-6 AgO3SCF3

SOL 75-09-2 CHZC12

CON 45 minutes, room temperature

\*\*STAGE(2)

RCT X 861404-04-8

CON SUBSTAGE(1) 17 hours, 45 deg C

SUBSTAGE(2) 45 deg C -> room temperature

\*\*STAGE(3)

SOL 64-17-5 EtOH

CON room temperature

PRO AG 861404-09-3

NTE in the dark

RX(15) RCT AG 861404-09-3

\*\*STAGE(1)

CAT 14221-01-3 Pd(PPh3)4

SOL 110-71-4 (CHZOMe)2

CON 30 minutes, room temperature

\*\*STAGE(2)

RCT AM 861404-10-6

RCT AJ 497-19-8 NAZCO3

SOL 7732-18-5 Water

CON SUBSTAGE(1) 4 hours, 80 deg C

SUBSTAGE(3)

SOL 7732-18-5 Water

CON room temperature

RX(35) OF 53 COMPOSED OF RX(13), RX(16)
RX(35) Y + X + AO ===> AP

Br

OMe OH

Whe Me

X

RX AO

RX

Me Me

MeO

OMe

Br

AP

AP
YIELD 68%

RX(13) RCT Y 18997-19-8

STAGE(1)
RGT 2 2923-28-6 Ag03SCF3
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature

STAGE (2)

L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

RCT X 861404-04-8

CON SUBSTAGE(1) 17 hours, 45 deg C

SUBSTAGE(3)

SOL 64-17-5 EtOH

CON room temperature

PRO AG 861404-09-3

NTE in the dark

RX(16) RCT AG 861404-09-3

STAGE(1)

CAT 14221-01-3 Pd(PPh3)4

SOL 110-71-4 (CH2OMe)2

CON 30 minutes, room temperature

STAGE(2)

RCT AO 23112-96-1

RGT AJ 497-19-8 NAZCO3

SOL 7732-18-5 Water

CON SUBSTAGE(3) 80 deg C

SUBSTAGE(3) 80 deg C -> room temperature

STAGE(3)

SOL 7732-18-5 Water

CON room temperature

PRO AP 861404-13-9

RX(49) OF 53 COMPOSED OF REACTION SEQUENCE RX(17), RX(15)

AND REACTION SEQUENCE RX(13), RX(15)

...AQ + AR ===> AM...

STEPS

AR

AQ

L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

L62 ANSWER 4 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

START NEXT REACTION SEQUENCE

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RX(17) RCT AQ 573-17-1

STAGE(1)

RGT AS 7553-56-2 I2, AT 75-03-6 EtI, AU 7439-95-4 Mg

SOL 109-99-9 THF

CON 1 hour, room temperature

) AV 121-43-7 He borate 109-99-9 THF SUBSTAGE(1) -78 deg C SUBSTAGE(2) -78 deg C -> room temperature STAGE (3) RCT AR 107-21-1 SOL 108-88-3 PhMe CON overnight, reflux PRO AM 861404-10-6 RX (13) RCT Y 18997-19-8 STAGE(1)
RGT Z 2923-28-6 AgO3SCF3
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature STAGE(2)
RCT X 861404-04-8
CON SUBSTAGE(1) 17 hours, 45 deg C
SUBSTAGE(2) 45 deg C -> room temperature STAGE(3) SOL 64-17-5 EtOH CON room temperature

(Continued)

PRO AG 861404-09-3 NTE in the dark

RX (15) RCT AG 861404-09-3

STAGE (2)

STAGE (1)

CAT 14221-01-3 Pd(PPh3)4
SOL 110-71-4 (CH2OMe)2
CON 30 minutes, room temperature

STAGE (2) AGE(2)
RCT AM 861404-10-6
RGT AJ 497-19-8 Na2CO3
SOL 7732-18-5 Water
CON SUBSTAGE(1) 4 hours, 80 deg C
SUBSTAGE(2) 80 deg C -> room temperature

STAGE(3) SOL 7732-18-5 Water CON room temperature PRO AN 861404-12-8

L62 ANSWER 5 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 143:60075 CASREACT
TITLE: Fixation of Both 02 and CO2 from Air by a Crystalline
Palladium Complex Bearing N-Heterocyclic Carbene Palladium Complex Bearing N-Heterocyclic Carbene Ligands Yamashita, Makoto: Goto, Kei; Kawashima, Takayuki Department of Chemistry, Graduate School of Science, University of Tokyo, Bunkyo, Tokyo, 113-0033, Japan Journal of the American Chemical Society (2005), 127(20), 7294-7295 COODN: JACSAT; ISSN: 0002-7863 American Chemical Society Journal English AUTHOR(S): CORPORATE SOURCE: PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI



Crystals of the two-coordinate Pd(0) complex Pd(ITmt)2 (1) bearing the

O2

and CO2 from air to produce the corresponding Pd(II) peroxocarbonate complex [Pd(ITmt)202002] (2). The present reaction consists of dioxygenation of the Pd(O) complex 1 to the Pd(II) peroxo complex [Pd(ITmt)202002] (3) and the subsequent CO2 insertion to produce the peroxocarbonate complex 2. Reaction of the Pd(O) peroxocarbonate complex 2. Reaction of the crystals of 1 with air was monitored by microscopic IR spectroscopy to confirm the sequence of the two-step solid-state reaction. The unique reactivity of solid 1 toward air was explained in terms of the structural features of the carbene ligand, ITmt.

REFERENCE COUNT:

32 THERE ARE 32 CITED REFERENCE

RECORD. ALL CITATIONS AVAILABLE IN THE RE

RX (4) OF 36 ...Q + N ===> R...

FORMAT

н2с≕ о a (4) → N

L62 ANSWER 5 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RX (4) RCT Q 50-00-0

STAGE (1)

AGE(1)
SOL 108-88-3 PhMe
CON SUBSTAGE(1) 120 deg C
SUBSTAGE(2) 120 deg C -> room temperature

STAGE (2) AGE(2)
RCT N 854030-35-6
RGT S 7647-01-0 HC1
SOL 60-29-7 Et20, 108-88-3 PhMe
CON SUBSTAGE(1) 1 hour, 120 deg C
SUBSTAGE(2) 120 deg C -> room temperature

PRO R 854030-33-4 NTE paraformaldehyde used

RX(13) OF 36 COMPOSED OF RX(4), RX(5) RX(13) Q + N ===> U

O H Q

L62 ANSWER 5 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

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STEPS
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\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RX (4) RCT Q 50-00-0 STAGE (1) GGE(1) SOL 108-88-3 PhMe CON SUBSTAGE(1) 120 deg C SUBSTAGE(2) 120 deg C -> room temperature STAGE(2)

RCT N 854030-35-6

RGT S 7647-01-0 HC1

SOL 60-29-7 ELCO, 108-88-3 PhMe

CON SUBSTAGE(1) 1 hour, 120 deg C

SUBSTAGE(2) 120 deg C -> room temperature PRO R 854030-33-4 NTE paraformaldehyde used R 854030-33-4 V 865-47-4 t-BUOK U 854030-37-9 60-29-7 Et20, 109-99-9 THF 1.5 hours, room temperature RX (5)

L62 ANSWER 6 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:

142:354850 CASREACT
Regarding the Mechanism of Olefin Metathesis with
Sol-Gel-Supported Ru-Based Complexes Bearing a
Bidentate Carbene Ligand. Spectroscopic Evidence for
Return of the Propagating Ru Carbene
Kingsbury, Jason S.; Hoveyda, Amir H.
Department of Chemistry, Merkert Chemistry Center,
Boston College, Chestnut Hill, MA, 02467, USA
Journal of the American Chemical Society (2005),
127(12), 4510-4517
CODEN: JACSAT: ISSN: 0002-7863
American Chemical Society
DOCUMENT TYPE:
Journal
LANGUAGE:
English
AB Two isotopically and structurally labeled Ru-based carbenes have been
prepared and attached to the surface of monolithic sol-gel glass. The
resulting glass-supported complexes exhibit significant catalytic
activity
In promoting olefin metathesis reactions and provide products of high
purity. Through anal. of the derivatized glass pellets used in a
sequence
of catalytic ring-closing metathesis reactions mediated by various
supported Ru carbenes, it is demonstrated that free Ru carbene
intermediates in solution can be scavenged by support-bound styrene ether
ligands prior to the onset of competing transition metal decomposition
The
observations detailed herein provide rigorous evidence that the initially

observations detailed herein provide rigorous evidence that the initially proposed release/return mechanism is, at least partially, operative. The present investigations shed light on a critical aspect of the mechanism

of an important class of Ru-based metathesis complexes (those bearing a bidentate styrene ether ligand).

REFERENCE COUNT: 57 THERE ARE 57 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

RX(16) OF 39 COMPOSED OF RX(2), RX(3) RX(16) C + J \*\*\*> K

L62 ANSWER 6 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

K: CM 1 YIELD 90%

K: CM 2 YIELD 90%

RX (2) RCT C 49673-43-0 STAGE(1)
RGT G 25895-60-7 NaBH3CN
SOL 67-56-1 MeOH
CON SUBSTAGE(1) room temperature -> 0 deg C
SUBSTAGE(2) 10 minutes, 0 deg C STAGE(2)

RGT H 7647-01-0 HCl

SOL 7732-18-5 Water

CON SUBSTAGE(1) 0 deg C, acidify

SUBSTAGE(2) 0 deg C -> 22 deg C

SUBSTAGE(3) 30 minutes, 22 deg C STAGE (3) RGT I 1310-58-3 KOH SOL 7732-18-5 Water CON room temperature, pH 8 - 9 PRO F 72991-60-7 NTE acidification in stage 2 repeated 3 times total F 72991-60-7, J 122-51-0 L 13826-83-0 NH4.BF4 K 848979-23-7 SUBSTAGE(1) room temperature SUBSTAGE(2) 10 hours, 120 deg C RX (3)

L62 ANSWER 6 OF 35 CASREACT COPYRIGHT 2006 ACS on STN SUBSTAGE(3) 120 deg C  $\rightarrow$  22 deg C (Continued)

L62 ANSWER 7 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

TITLE:

\$142:347439 CASREACT

Synthesis, spectroscopic and electrochemical properties of some heteroleptic tris-chelates of ruthenium(II) involving 2,2'-bipyridine (bpy) and N-(aryl)pyridine-2-aldimine (L): X-ray crystal atructures of [Ru(bpy)(L2)2](Cl04)2-H2O and 3-N-(4-tolyl)imidazo(1,5a]pyridinium perchlorate Hishra, Dipankar; Naskar, Subhendu; Adhikary, Bibhutosh; Butcher, Raymond J.; Chattopadhyay,

L3 = (4-chlorophenyl)(2-pyridylmethylene)amine, L4 = (4-fluorophenyl)(2-pyridylmethylene)amine, D4 = (4-chlorophenyl)(2-pyridylmethylene)amine and bpy = 2,2'-bipyridyl) were synthesized. In addition to these ruthenium complexes, the authors also were able to isolate four imidazopyridnium perchlorate componed. B1-B4 from the same reactions. The x-ray crystal structures of one representative ruthenium complex (A2) and the imidazopyridnium perchlorate compound (B2) were determined The Ru(II) center in the complex is coordinated by six N donors with a distorted octahedral geometry. The imine ligands (L) act as bidentate N,N donors. REFERENCE COUNT: THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

A + 3 B + C ==> D + E RX(1) OF 4

L62 ANSWER 7 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RX(2) RCT A 69141-04-4, G 7471-13-8, C 67-56-1

STAGE(1)
SOL 67-56-1 MeON
CON SUBSTAGE(1) room temperature -> reflux
SUBSTAGE(2) 4 hours, reflux
SUBSTAGE(3) reflux -> room temperature

RGT F 7601-89-0 NaClO4 SOL 67-56-1 MeOH CON room temperature

PRO H 156843-38-8, I 738585-87-0 NTE safety - product is a potentially explosive perchlorate salt

L62 ANSWER 7 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

$$H_3C^{-s-OH}$$

$$C \qquad \xrightarrow{(1)} \qquad O = \begin{bmatrix} 0 \\ C1 \\ O \end{bmatrix} - O^-$$

$$D: CM 1$$

RCT A 69141-04-4, B 7032-25-9, C 67-56-1 RX (1)

STAGE(1)
SOL 67-56-1 MeOH
SOL SUBSTAGE(1) room temperature -> reflux
SUBSTAGE(2) 4 hours, reflux
SUBSTAGE(3) reflux -> room temperature

STAGE (2)
RGT F 7601-89-0 NaClO4
SOL 67-56-1 MeOH
CON room temperature

PRO D 848303-98-0, E 848304-04-1 NTE safety - product is a potentially explosive perchlorate salt

RX (2) OF 4 A + 3 G + C ===> H + I

L62 ANSWER 7 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

RX(3) OF 4 A + 3 J + C ===> K + L

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RX (3) RCT A 69141-04-4, J 26825-34-3, C 67-56-1

STAGE(1)
SOL 67-56-1 MeOH
CON SUBSTAGE(1) room temperature -> reflux
SUBSTAGE(2) 4 hours, reflux
SUBSTAGE(3) reflux -> room temperature

STAGE(2)

RGT F 7601-89-0 NaClO4

SOL 67-56-1 MeOH

CON room temperature

L62 ANSWER 7 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)
PRO K 848304-00-7, L 738585-89-2
NTE safety - product is a potentially explosive perchlorate salt

RX (4) OF 4 A + 3 N + C ===> N + O

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RX (4) RCT A 69141-04-4, M 29202-06-0, C 67-56-1

STAGE (1)

SOL 67-56-1 MeOH
CON SUBSTAGE(1) room temperature -> reflux
SUBSTAGE(2) 4 hours, reflux
SUBSTAGE(3) reflux -> room temperature

L62 ANSWER 8 OF 35
ACCESSION NUMBER:
142:93738 CASREACT
TITLE:
Sterically demanding, bioxazoline-derived
N-heterocyclic carbene ligands with restricted
flexibility for catalysis
Altenhoff, Gereon; Goddard, Richard; Lehmann,
Christian W.; Glorius, Frank
MAX-Planck-Institut fuer Kohlenforschung, Muelheim an
der Ruhr, 45470, Germany
Journal of the American Chemical Society (2004),
126(46), 15195-15201
CODEN: JACSAT; ISSN: 0002-7863
American Chemical Society
Journal

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

The triflate salts of imidazobioxazolium ions I [R = Rl = Me; RRl = [CR2]n: n = 5, 6, 7, 8, 12] are prepared as precursors for sterically demanding and conformationally constrained N-heterocyclic carbene (NNC) ligands; palladium complexes derived from I [RRl = (CR2)n: n = 7, 12] act as effective catalysts for the Suzuki-Miyaura coupling reactions of ortho-substituted aryll chlorides with ortho-substituted aryllopronic acids to provide triortho- and tetraortho-substituted blaryla such as II in 47-961 yields. I=cr3503-are prepared in five steps from a,a-disubstituted amino acids and di-Et oxalate: reduction of amino acids to the amino alcs., condensation of the amino alcs. with

di-Et

oxalate to give the hydroxymethyl-substituted oxamides, chlorination of
the primary alc. moleties, cyclization of the oxamide with the
chloromethyl groups to give the bioxazolines, and reaction of the
bioxazolines with chloromethyl pivalet and silver triflate.
I=CF3503- are soluble in methylene chloride and THF and are
chromatographable. Iridium cyclooctadienyl and iridium dicarbonyl
chloride complexes derived from I=CF3503- [R = Rl = Me; RRl = (CH2)n;
n = 6, 8, 12] are prepared; IR frequencies of the carbonyl ligands
indicate

that carbene ligands derived from I-CF3503- are less electron-donating than previous NHC ligands but are comparable to electron-rich phosphines. Selected iridium cyclooctadienyl and iridium dicarbonyl chloride

lexes
of imidazobioxazolium ligands are characterized by X-ray crystallog.
Dimeric palladium chloride complexes derived from I=CF3SO3- [RR] =
(CH2)n; n = 7, 12] are prepared and characterized by X-ray crystallog.
Generation of the carbene ligand from I=CF3SO3- [RR] = (CH2)12] by
treatment with potassium hydride and potassium tert-butoxide followed by
addition of palladium acetate yields a palladium catalyst which is
tive

for the Suzuki-Miyaura coupling of highly hindered aryl chlorides and

L62 ANSWER 7 OF 35 CASREACT COPYRIGHT 2006 ACS on STN STAGE(2) (Continued)

) F 7601-89-0 NaClO4 67-56-1 MeOH room temperature RGT SOL CON

N 848304-02-9, O 848304-06-3 safety - product is a potentially explosive perchlorate salt

L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) arylboronic acids. Potassium phosphate is the most effective base and toluene is the most effective solvent for Suzuki-Miyaura coupling of highly hindered aryl chlorides and arylboronic acids using imidazobioxazolium-derived carbens ligands, although cesium carbonate can also be used as the base and 1,4-dioxane as the solvent; the isolated dimeric palladium chloride complexes derived from !ccf3303-[RR] = (CH2)n; n = 7, 12] can also be used as catalysts. Anhyd. conditions are important to minimize hydrodeborylation byproducts of the coupling reaction. E.g., in the presence of the palladium catalyst generated from !ccf3303-[RR] = (CH2)12] and palladium acctate and potassium phosphate, 2-chloro-1,3-dimethylbenzene and 2,4,6-trimethylphenylboronic acid undergo coupling in toluene at 100° for 16 h to provide biphenyl II in 968 yield.

REFERENCE COUNT: 67 THERE ARE 67 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

RX(23) OF 154 ...AP + AQ + AI ===> AR...

RCT AP 18997-19-8, AO 2923-28-6 RX (23)

STAGE (1)

SOL 75-09-2 CH2C12 CON 45 minutes, room temperature

STAGE (2)

AI 49585-66-2 75-09-2 CH2C12 SUBSTAGE(1) 20 hours, 40 deg C SUBSTAGE(2) 40 deg C -> room temperature

STAGE (3)

```
L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN RGT E 67-56-1 MeOH
                                                                                                                             L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN STAGE(3) RGT E 67-56-1 HeOH
                                                                                     (Continued)
                                                                                                                                                                                                                  (Continued)
              PRO AR 814254-77-8
NTE sealed tube (2nd stage), in the dark (2nd stage)
                                                                                                                                           PRO AT 814254-79-0 NTE sealed tube (2nd stage), in the dark (2nd stage)
RX(24) OF 154 ...AP + AQ + AK ===> AT
                                                                                                                              RX(25) OF 154 ...AP + AQ + AL ***> AU...
                                                                                                                                                                                                 (25)
                                                                                                                              AU: CM 1
YIELD 85%
AT: CM 2
YIELD 65%
                                                                                                                                                     AU: CM 2
YIELD 85%
RX (24)
          RCT AP 18997-19-8, AQ 2923-28-6
                                                                                                                              RX (25)
                                                                                                                                          RCT AP 18997-19-8, AQ 2923-28-6
                STAGE(1)
SOL 75-09-2 CH2Cl2
CON 45 minutes, room temperature
                                                                                                                                              STAGE(1)
SOL 75-09-2 CH2Cl2
CON 45 minutes, room temperature
               STAGE(2)
RCT AK 814254-72-3
SOL 75-09-2 CH2C12
CON SUBSTAGE(1) 20 hours, 40 deg C
SUBSTAGE(2) 40 deg C -> room temperature
                                                                                                                                             STAGE(2)

RCT AL 606970-67-6

SOL 75-09-2 CH2C12

CON SUBSTAGE(1) 20 hours, 40 deg C

SUBSTAGE(2) 40 deg C -> room temperature
                                                                                                                             L62 ANSWER 8 0 0F 35 CASREACT COPYRIGHT 2006 ACS on STN 50L 75-09-2 CH2Cl2 CON SUBSTAGE(1) 20 hours, 40 deg C SUBSTAGE(2) 40 deg C -> room temperature
L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                  (Continued)
                STAGE(3)
RGT E 67-56-1 MeOH
             PRO AU 606970-69-8
NTE sealed tube (2nd stage), in the dark (2nd stage)
                                                                                                                                              STAGE(3)
RGT E 67~56-1 MeOH
                                                                                                                                           PRO AV 814254-81-4
NTE sealed tube (2nd stage), in the dark (2nd stage)
                     ...AP + AQ + AN ===> AV...
RX (26) OF 154
                                                                                                                                                     ...AP + AQ + AN ===> AW...
                                                                                                                                                            ● Ag(I)
   ● Ag(I)
                                                                       (26)
                                                                                                                                                                                             AW: CM 1
YIELD 61%
                       AV: CM 2
YIELD 63%
             RCT AP 18997-19-8, AQ 2923-28-6
RX (26)
                STAGE(1)
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature
                                                                                                                              AW: CM 2
YIELD 61%
```

RX(27) RCT AP 18997-19-8, AQ 2923-28-6

STAGE (2) RCT AM 814254-73-4 L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

STAGE(1)

SOL 75-09-2 CH2C12

CON 45 minutes, room temperature

STAGE(2)

RCT AN 814254-74-5

SOL 75-09-2 CH2C12

CON SUBSTAGE(1) 20 hours, 40 deg C

SUBSTAGE(2) 40 deg C -> room temperature

STAGE(3)

RGT E 67-56-1 MeOH

PRO AN 814254-83-6

NTE sealed tube (2nd stage), in the dark (2nd stage)

RX(28) OF 154 ...AP + AQ + AO ===> AX...

AP

Ag(1)

AQ

AO

(28)

● Ag(I)

L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

F-C-SO3-

RX(69) OF 154 COMPOSED OF RX(17), RX(23) RX(69) AA + AP + AQ ===> AR

L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

2
STEPS

F-C-SO3
Me Me Me

AR: CM 1
YIELD 83%

RX (17)

RCT AA 61051-14-7

RGT F 1310-73-2 NaOH
PRO AI 49585-66-2

SOL 64-17-5 EtcH, 109-99-9 THF
CON SUBSTAGE(1) 30 minutes, room temperature

SUBSTAGE(2) 3 hours, 90 deg C

RX (23)

RCT AP 18997-19-8, AQ 2923-28-6

STAGE (1)

SOL 75-09-2 CH2C12

CON 45 minutes, room temperature

STAGE (2)

RCT AI 49585-66-2

SOL 75-09-2 CH2C12

CON 45 SUBSTAGE(1) 20 hours, 40 deg C

SUBSTAGE(2) 40 deg C -> room temperature

STAGE (3)

RGT E 67-56-1 MeOH

PRO AR 814254-77-8

NTE sealed tube (2nd stage), in the dark (2nd stage)

RX (70) OF 154 COMPOSED OF RX (18), RX (24)
RX (70) AD + AP + AQ ===> AT

AP

```
L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                          (Continued)
                                                                                                                                                            L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                                                                                                                                                                                       (Continued)
STEPS
                    AU: CM 1
YIELD 85%
                                                                                                                                                                                AV: CM 1
YIELD 63%
                                                 AU: CM 2
YIELD 85%
                                                                                                                                                                                                             AV: CM 2
YIELD 63%
                 RCT AE 606970-66-5
RGT F 1310-73-2 NaOH
PRO AL 606970-67-6
SOL 64-17-5 EtOH, 109-99-9 THF
CON SUBSTAGE(1) 30 minutes, room temperature
SUBSTAGE(2) 3 hours, 90 deg C
RX (19)
                                                                                                                                                                             RCT AF 814254-69-9
RGT F 1310-73-2 NaOH
PRO AM 814254-73-4
SOL 64-17-5 EtOH, 109-99-9 THF
CON SUBSTAGE(1) 30 minutes, room temperature
SUBSTAGE(2) 3 hours, 90 deg C
                                                                                                                                                            RX (20)
RX (25)
                 RCT AP 18997-19-8, AQ 2923-28-6
                    STAGE(1)
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature
                                                                                                                                                            RX (26)
                                                                                                                                                                             RCT AP 18997-19-8, AQ 2923-28-6
                                                                                                                                                                                STAGE(1)
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature
                   STAGE(2)

RCT AL 606970-67-6

SOL 75-09-2 CH2C12

CON SUBSTAGE(1) 20 hours, 40 deg C

SUBSTAGE(2) 40 deg C -> room temperature
                                                                                                                                                                                STAGE(2)

RCT AM 814254-73-4

SOL 75-09-2 CH2C12

CON SUBSTAGE(1) 20 hours, 40 deg C

SUBSTAGE(2) 40 deg C -> room temperature
                                                                                                                                                                                 STAGE (3)
RGT E 67-56-1 MeOH
                 PRO AU 606970-69-8
NTE sealed tube (2nd stage), in the dark (2nd stage)
                                                                                                                                                                             PRO AV 814254-81-6
NTE sealed tube (2nd stage), in the dark (2nd stage)
RX(72) OF 154 COMPOSED OF RX(20), RX(26)

RX(72) AF + AP + AQ ===> AV
                                                                                                                                                           RX (73) OF 154 COMPOSED OF RX (21), RX (27)
RX (73) AG + AP + AQ ===> AW
                                                           AP
                                                                                               ΑQ
                                                                                                                                                           L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN SOL 75-09-2 CH2C12 CON SUBSTAGE(1) 20 hours, 40 deg C SUBSTAGE(2) 40 deg C -> room temperature
L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                          (Continued)
                                                                                                                                                                                                                                                                      (Continued)
                                                                                                                                                                                STAGE(3)
RGT E 67-56-1 MeOH
                                                                                                                                                                              PRO AW 814254-83-6
NTE sealed tube (2nd stage), in the dark (2nd stage)
                                                              c1<sup>-1</sup>
                                                              AP
                                                                                                                                                            RX (74) OF 154 COMPOSED OF RX (22), RX (28) RX (74) AH + AP + AQ ===> AX
                                                                                                                                                                                                                                                 cı ~ o
                            STEPS
                                                                                                                                                                                                                                                 AP
                                                                                                                                                                                        STEPS
                                                                                                                                                                                                            AX: CM 1
YIELD 60%
AW: CM 2
YIELD 61%
                       AG 814254-70-1
F 1310-73-2 NaOH
AN 814254-74-5
64-17-5 EtCH, 109-99-9 THF
SUBSTAGE(1) 30 minutes, room temperature
SUBSTAGE(2) 3 hours, 90 deg C
RX(21)
RX (27)
                 RCT AP 16997-19-8, AQ 2923-28-6
                    STAGE(1)
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature
                                                                                                                                                            AX: CM 2
YIELD 60%
```

STAGE (2) RCT AN 814254-74-5

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L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS ON STN RX(22) RCT AH 914254-71-2 RGT F 1310-73-2 NaOH PRO AO 814254-75-6 SOL 64-17-5 EtOH, 109-99-9 THF SUBSTAGE(1) 30 minutes, room temperature SUBSTAGE(2) 3 hours, 90 deg C
                                                                                                                (Continued)
                                                                                                                                                                    L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN RX(11) RCT R 61051-10-3
                                                                                                                                                                                                                                                                                     (Continued)
                                                                                                                                                                                          STAGE (1)
                                                                                                                                                                                               AGE(1)
RGT AB 7719-09-7 SOC12
SOL 108-88-3 PhMe
CON SUBSTAGE(1) 1 hour, 60 deg C
SUBSTAGE(2) 3 hours, 90 deg C
SUBSTAGE(3) 90 deg C -> room temperature
                  RCT AP 18997-19-8, AQ 2923-28-6
RX (28)
                     STAGE(1)
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature
                                                                                                                                                                                          STAGE (2)
                                                                                                                                                                                                RGT AC 1310-58-3 KOH
SOL 7732-18-5 Water
                     STAGE (2)
                                                                                                                                                                                       PRO AA 61051-14-7
                           MGE(2)

RCT AO 814254-75-6

SOL 75-09-2 CHZC12

CON SUBSTAGE(1) 20 hours, 40 deg C

SUBSTAGE(2) 40 deg C -> room temperature
                                                                                                                                                                                             AA 61051-14-7
F 1310-73-2 NAOH
AI 49585-66-2
64-17-5 ELOH, 109-99-9 THF
SUBSTAGE(1) 30 minutes, room temperature
SUBSTAGE(2) 3 hours, 90 deg C
                                                                                                                                                                    RX (17)
                                                                                                                                                                                       RCT
                                                                                                                                                                                       PRO
                     STAGE (3)
RGT E 67-56-1 MeOH
                  PRO AX 814254-85-8
NTE sealed tube (2nd stage), in the dark (2nd stage)
                                                                                                                                                                    RX (23)
                                                                                                                                                                                      RCT AP 18997-19-8, AQ 2923-28-6
                                                                                                                                                                                          STAGE(1)
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature
RX(99) OF 154 COMPOSED OF RX(11), RX(17), RX(23)
RX(99) R + AP + AQ ===> AR
                                                                                                                                                                                         STAGE(2)
RCT AI 49585-66-2
SOL 75-09-2 CH2C12
CON SUBSTAGE(1) 20 hours, 40 deg C
SUBSTAGE(2) 40 deg C -> room temperature
                                                                                                                                                                                          STAGE(3)
RGT E 67-56-1 MeOH
                                                               AP
                                                                                                        ● Ag(I)
                                                                                                                                                                                       PRO AR 814254-77-8
NTE sealed tube (2nd stage), in the dark (2nd stage)
                                                                                                    AO
                                                                                                                                                                    RX(101) OF 154 COMPOSED OF RX(12), RX(18), RX(24) RX(101) T + AP + AQ ===> AT
STEPS
                                                    AR: CM 2
YIELD 83%
                                                                                                                                                                                                                                                                      ● Ag(I)
                                                                                                                                                                                                                                                                  AO
L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                                (Continued)
                                                                                                                                                                    L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                                                                                                                                                                                                    .CF3
STEPS
                                                                                                                                                                                                                              cı'
                                                                                                                                                                                                                                                                        • Ag(I)
                                                                                                                                                                                                                              ΑP
                                                   AT: CM 2
YIELD 65%
                                                                                                                                                                    STEPS
RX (12)
                  RCT T 814254-64-3
                    STAGE(1)
RGT AB 7719-09-7 SOC12
SOL 108-88-3 PhMe
CON SUBSTAGE(1) 1 hour, 60 deg C
SUBSTAGE(2) 3 hours, 90 deg C
SUBSTAGE(3) 90 deg C -> room temperature
                                                                                                                                                                    r-c-so3
                     STAGE(2)
RGT AC 1310-58-3 KOH
SOL 7732-18-5 Water
                                                                                                                                                                    AU: CM 1
YIELD 85%
                                                                                                                                                                                                   AU: CM 2
YIELD 85%
                  PRO AD 814254-68-7
                 RCT AD 814254-68-7

RGT F 1310-73-2 NaOH

PRO AM 814254-72-3

SOL 64-17-5 ECOH, 109-99-9 THF

CON SUBSTAGE(1) 30 minutes, room temperature

SUBSTAGE(2) 3 hours, 90 deg C
RX (18)
                                                                                                                                                                    RX(13)
                                                                                                                                                                                      RCT V 101725-44-4
                                                                                                                                                                                        STAGE(1)

RGT AB 7719-09-7 SOC12

SOL 108-88-3 PhMe

CON SUBSTAGE(1) 1 hour, 60 deg C

SUBSTAGE(2) 3 hours, 90 deg C

SUBSTAGE(3) 90 deg C -> room temperature
RX (24)
                 RCT AP 18997-19-8, AQ 2923-28-6
                     STAGE (1)
                          SOL 75-09-2 CH2Cl2
CON 45 minutes, room temperature
                                                                                                                                                                                                RGT AC 1310-58-3 KOH
SOL 7732-18-5 Water
                     STAGE (2)
                           AGE (2)

RCT AK 814254-72-3

SOL 75-09-2 CHZC12

CON SUBSTAGE (1) 20 hours, 40 deg C

SUBSTAGE (2) 40 deg C -> room temperature
                                                                                                                                                                                       PRO AE 606970-66-5
                                                                                                                                                                                      RCT AE 606970-66-5
RGT F 1310-73-2 NaOH
PRO AL 606970-67-6
SOL 64-17-5 EtOH, 109-99-9 THF
CON SUBSTAGE(1) 30 minutes, room temperature
SUBSTAGE(2) 3 hours, 90 deg C
                                                                                                                                                                    RX (19)
                     STAGE (3)
RGT E 67-56-1 MeOH
                  PRO AT 914254-79-0
NTE sealed tube (2nd stage), in the dark (2nd stage)
                                                                                                                                                                    RX (25)
                                                                                                                                                                                      RCT AP 18997-19-8, AQ 2923-28-6
                                                                                                                                                                                          STAGE(1)
SOL 75-09-2 CH2Cl2
CON 45 minutes, room temperature
RX(103) OF 154 COMPOSED OF RX(13), RX(19), RX(25) RX(103) V + AP + AQ ===> AU
```

L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN SOL 7732-18-5 Water (Continued) STAGE(2)
RCT AL 606970-67-6
SOL 75-09-2 CH2C12
CON SUBSTAGE(1) 20 hours, 40 deg C
SUBSTAGE(2) 40 deg C -> room temperature PRO AF 814254-69-8 RCT AF 814254-69-8
RGT F 1310-73-2 NAOH
PRO AM 814254-73-4
SOL 64-17-5 EtOH, 109-99-9 THF
CON SUBSTAGE(1) 30 minutes, room temperature
SUBSTAGE(2) 3 hours, 90 deg C RX (20) STAGE (3) RGT E 67-56-1 MeOH PRO AU 606970-69-8 NTE sealed tube (2nd stage), in the dark (2nd stage) RX (26) RCT AP 18997-19-8, AQ 2923-28-6 STAGE(1) SOL 75-09-2 CH2C12 CON 45 minutes, room temperature RX(105) OF 154 COMPOSED OF RX(14), RX(20), RX(26)RX(105) X + AP + AQ ===> AV STAGE(2)
RCT AM 814254-73-4
SOL 75-09-2 CH2C12
CON SUBSTAGE(1) 20 hours, 40 deg C
SUBSTAGE(2) 40 deg C -> room temperature STAGE (3) RGT E 67+56-1 MeOH cı′ ● Aq(I) PRO AV 814254-81-4 NTE sealed tube (2nd stage), in the dark (2nd stage) ΑP ΑO RX(107) OF 154 COMPOSED OF RX(15), RX(21), RX(27) RX(107)  $\tau$  + AP + AQ ===> AW STEPS AV: CM 1 YIELD 63% AV: CM 2 YIELD 63% C1 RX (14) RCT X 814254-65-4 STAGE(1)

RGT AB 7719-09-7 SOC12

SOL 108-88-3 PhMe

CON SUBSTAGE(1) 1 hour, 60 deg C

SUBSTAGE(2) 3 hours, 90 deg C

SUBSTAGE(3) 90 deg C -> room temperature STAGE(2) RGT AC 1310-58-3 KOH L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
RCT AN 814254-74-5
SOL 75-09-2 CH2C12
CON SUBSTAGE(2) 40 deg C -> room temperature L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) STAGE (3) RGT E 67-56-1 MeOH PRO AW **814254-83-6** NTE sealed tube (2nd stage), in the dark (2nd stage) ● Ag(I) STEPS AW: CM 1 YIELD 61% RX(109) OF 154 COMPOSED OF RX(16), RX(22), RX(28) RX(109) E + AP + AQ ===> AX AW: CM 2 YIELD 61% RX (15) RCT Y 814254-66-5 STAGE(1)

RGT AB 7719-09-7 SOC12

SOL 108-88-3 PhMe

CON SUBSTAGE(1) 1 hour, 60 deg C

SUBSTAGE(2) 3 hours, 90 deg C

SUBSTAGE(3) 90 deg C -> room temperature ● Aq(I) STEPS AX: CH 1 YIELD 60% STAGE(2) RGT AC 1310-58-3 KOH SOL 7732-18-5 Water PRO AG 814254-70-1 RCT AG 814254-70-1 RGT F 1310-73-2 NaOH PRO AN 814254-74-5 SOL 64-17-5 EtOH, 109-99-9 THF SOL SUBSTAGE[1] 30 minutes, room temperature SUBSTAGE[2] 3 hours, 90 deg C RX (21) RX (27) RCT AP 18997-19-8, AQ 2923-28-6 STAGE(1)
SOL 75-09-2 CH2C12
CON 45 minutes, room temperature AX: CM 2 YIELD 60%

STAGE (2)

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L62 ANSWER 8 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                                (Continued)
                 RCT Z 814254-67-6
RX (16)
                     STAGE (1)
                                   )
AB 7719-09-7 SOC12
108-88-3 PhMe
SUBSTAGE(1) 1 hour, 60 deg C
SUBSTAGE(2) 3 hours, 90 deg C
SUBSTAGE(3) 90 deg C -> room temperature
                           RGT
SOL
CON
                     STAGE (2)
                           RGT AC 1310-58-3 KOH
SOL 7732-18-5 Water
                  PRO AH 814254-71-2
                 RCT AH 814254-71-2
RGT F 1310-73-2 NAOH
PRO AO 814254-75-6
SOL 64-17-5 EtOH, 109-99-9 THF
CON SUBSTAGE(1) 30 minutes, room temperature
SUBSTAGE(2) 3 hours, 90 deg C
RX (22)
                 RCT AP 18997-19-8. AO 2923-28-6
RX (28)
                     STAGE (1)
                          SOL 75-09-2 CH2Cl2
CON 45 minutes, room temperature
                      STAGE (2)
                          AGE (1)

RCT AO 814254-75-6

SOL 75-09-2 CH2C12

CON SUBSTAGE(1) 20 hours, 40 deg C

SUBSTAGE(2) 40 deg C -> room temperature
                     STAGE (3)
RGT E 67-56-1 MeOH
```

PRO AX 814254-85-8 NTE sealed tube (2nd stage), in the dark (2nd stage)

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L62 ANSWER 9 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 141:395258 CASREACT
TITLE: A Benzimidazole-Based N-Heterocyclic Carbene Derived from 1,10-Phenanthroline
AUTHOR(S): Hetallinos, Costa; Barrett, Fred B.; Chaytor,
Jennifer
                                                            L.; Heska, Mary E. A.
Department of Chemistry, Brock University, St.
Catharines, ON, L2S 3A1, Can.
Organic Letters (2004), 6(20), 3641-3644
CODEN: ORLEF7; ISSN: 1523-7060
American Chemical Society
Journal
CORPORATE SOURCE:
SOURCE:
 PUBLISHER:
 DOCUMENT TYPE:
LANGUAGE:
          MUMGE: English
A Catalytically active palladium-complexed tetracyclic N-heterocyclic
carbene (NMC) was prepared in three steps from com. available
1,10-phenanthroline by using a reduction-cyclization-deprotonation
 sequence.
The new carbene framework is a prototype for the development of a series of chiral N-heterocyclic carbenes.
REFERENCE COUNT: 42 THERE ARE 42 CITED REFERENCES AVAILABLE FOR
REFERENCE COUNT:
                                                                          RECORD. ALL CITATIONS AVAILABLE IN THE RE
FORMAT
RX\{10\} OF 17 COMPOSED OF RX\{3\}, RX\{4\}

RX\{10\} A + J ===> K
                                                                                             STEPS
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L62 ANSWER 9 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) K YIELD 89% RX (3) RCT A 66-71-7 STAGE (1) AGE(1)
RGT C 25895-60-7 NABH3CN
SOL 67-56-1 MeOH, 64-19-7 AcOH
CON SUBSTAGE(1) 0.08 hours, room temperature
SUBSTAGE(2) room temperature -> reflux
SUBSTAGE(3) 6 hours, reflux
SUBSTAGE(4) reflux -> room temperature

RGT H 1310-73-2 NaOH SOL 7732-18-5 Water, 67-56-1 MeOH CON room temperature, pH 12

GGE(1) RGT L 7647-01-0 HC1 SOL 7732-18-5 Water, 122-51-0 CH(OEt)3 CON 15 hours, 80 deg C

RCT G 56798-33-5, J 122-51-0

STAGE(2) RGT M 7782-44-7 O2 CON 2 hours, 80 deg C

PRO G 56798-33-5
NTE optimization study, other products also detected, optimized on solvent, product depends on solvent, incremental addition of NaBH3CN

STAGE (2)

STAGE (1)

PRO K 786688-16-2 RX(15) OF 17 COMPOSED OF RX(3), RX(4), RX(8) RX(15) A + J + AA ===> AB

RX (4)

● Na<sup>1</sup> AA AB: CM 1 YIELD 90% AB: CM 2 YIELD 90% RCT A 66-71-7 RX (3) STAGE(1) AGE(1)
RGT (25895-60-7 NABH3CN
SOL 67-56-1 MeOH, 64-19-7 ACOH
CON SUBSTAGE(1) 0.08 hours, room temperature
SUBSTAGE(2) room temperature -> reflux
SUBSTAGE(3) 6 hours, reflux
SUBSTAGE(4) reflux -> room temperature STAGE(2) RGT H 1310-73-2 NaOH SOL 7732-18-5 Water, 67-56-1 MeOH CON room temperature, pH 12 PRO G 56798-33-5 NTE optimization study, other products also detected, optimized on

L62 ANSWER 9 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

(Continued)

STEPS

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L62 ANSWER 9 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) solvent, product depends on solvent, incremental addition of NaBHSCN
RX (4)
                  RCT G 56798-33-5, J 122-51-0
                    STAGE(1)

RGT L 7647-01-0 HC1

SOL 7732-18-5 Water, 122-51-0 CH(OEt)3

CON 15 hours, 80 deg C
                    STAGE(2)

RGT M 7782-44-7 O2

CON 2 hours, 80 deg C
                  PRO K 786688-16-2
                 RCT K 786688-16-2, AA 143-66-8
PRO AB 786688-19-5
SOL 67-56-1 MeOH
CON room temperature
RX (8)
```

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L62 ANSWER 10 OF 35

ACCESSION NUMBER:
141:140590 CASREACT
141:140590 CASREACT
TITLE:
A New Class of Chelating N-Heterocyclic Carbene
Ligands and Their Complexes with Palladium

AUTHOR(3):
Waltman, Andrew W., Grubbs, Robert H.

Division of Chemistry and Chemical Engineering,
California Institute of Technology, Pasadena, CA,
91125, USA

SOURCE:
Organometallics (2004), 23(13), 3105-3107

CODEN: ORGND7; ISSN: 0276-7333

PUBLISHER:
American Chemical Society
DOCUMENT TYPE:
Journal
LANGUAGE:
English
AB A new series of chelating N-(o-phenolato)-N-heterocyclic carbene (NHC)
ligands mimicking salicylaldimine framework and their palladium complexes
are described. General synthetic pathway to N-hydroxyaryl-substituted
imidazolidinylidenes is described, starting from unsym. oxalodiamy oxalodiamies
ARINCOCONHAR1, (2a-d) where Ar = 2,4,6-Me306H2 or 2,6-iPr2C6H3 (Mes and
Dipp, resp.) and ArlH = 2-H0-3-R1-5-R2C6H2 (R1, R2 = H; EBU, Me;
1-adamantyl, Me). Reduction of 2a-d followed by condensation with
orthoformate gave imidazolium salts 1-Arl-3-Ar-4,5-dihydro-1H-imidazolium
chlorides (4a-d), which were converted to potassium carbenes-phenolates
[ARINCHGNAR1]-K- and reacted with palladium dimers [Pd(PR3)MeC1]? to give
[(PR3)PdMe[arx0H80Arl-4C2,XO]] (8a-d, PR3 = PR3, PP3), PEC1).
Crystel structures of complexes 8a (R1 = R2 = H; PR3 = PEC3) and 8d (R1 =
1-adamantyl, R2 Mer PR3 = PPR3) are reported. The ligands feature a
chelating phenolic unit, thereby expanding the class of available NHC
ligands for organometallic catalysis.

REFERENCE COUNT:
38 THERE ARE 38 CITED REFERENCES AVAILABLE FOR
THIS
                                                                                                                                                                                                                                                                                                                         RECORD. ALL CITATIONS AVAILABLE IN THE RE
         FORMAT
            RX(11) OF 85 ...I + AG ===> AH...
```

ELO AG

L62 ANSWER 10 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) AH YIELD 55% RX (11) RCT I 724794-61-0 STAGE (1) RGT AI 14044-65-6 BH3-THF
SOL 109-99-9 THF
CON SUBSTAGE(1) overnight, reflux
SUBSTAGE(2) reflux -> room temperature STAGE (2) RGT T 67-56-1 MeOH STAGE (3) RGT N 7647-01-0 HCl SOL 7732-18-5 Water STAGE (4)

RCT AG 122-51-0

SOL 122-51-0 CH(OEt) 3

CON SUBSTAGE(1) room temperature -> 100 deg C

SUBSTAGE(2) 6 minutes, 100 deg C PRO AH 724794-66-5 RX(12) OF 85 ...K + AG ===> AJ...

L62 ANSWER 10 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) (12) AJ YIELD 85% RX(12) RCT K 724794-62-1 STAGE(1)

RGT AI 14044-65-6 BH3-THF

SOL 109-99-9 THF

CON SUBSTAGE(1) overnight, reflux

SUBSTAGE(2) reflux -> room temperature STAGE(2) RGT T 67-56-1 MeOH STAGE(3) RGT N 7647-01-0 HCl SOL 7732-18-5 Water STAGE(4)

RCT AG 122-51-0

SOL 122-51-0 CH(OEt)3

CON SUBSTAGE(1) room temperature -> 100 deg C

SUBSTAGE(2) 6 minutes, 100 deg C

L62 ANSWER 10 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) PRO AJ 724794-67-6

RX (13) OF 85 ...X + AG ===> AR...

• c1 AK YIELD 35%

RX (13) RCT X 724794-63-2

STAGE (1)

, AI 14044-65-6 BH3-THF 109-99-9 THF SUBSTAGE(1) overnight, reflux SUBSTAGE(2) reflux -> room temperature

L62 ANSWER 10 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

STAGE(2) RGT T 67-56-1 MeOH

STAGE (3) RGT N 7647-01-0 HC1 SOL 7732-18-5 Water

STAGE(4)

RCT AG 122-51-0

SOL 122-51-0 CH(OEt)3

CON SUBSTAGE(1) room temperature -> 100 deg C

SUBSTAGE(2) 6 minutes, 100 deg C

PRO AK 724794-68-7

L62 ANSWER 11 OF 35
ACCESSION NUMBER:
TITLE:
Preparation of axially chiral N,N'-diarylimidazolium and N-arylthiazolium salts and evaluation of their catalytic potential in the benzoin and in the intramolecular Stetter reactions

AUTHOR(S):
CORPORATE SOURCE:
SOURCE:
SOURCE:
Pesch, Jens; Narms, Klaus; Bach, Thoraten
Lehrstuhl Zuer Organische Chemie I, Technische
Universitaet Muenchen, Garching, 85747, Germany
European Journal of Organic Chemistry (2004), (9),
2023-2035
COBDE: EJOCFK; ISSN: 1434-193X
Wiley-VCH Verlag GmbH & Co. KGAA
Journal
LANGUAGE:
English

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

AB N-Aryl-substituted imidazoles were prepared which contain a stereogenic axis

and which can occur as atropisomers. The
di(2-1sopropylphenyl) imidazolium
salts could be obtained from 2-isopropylaniline and diacetyl in three
steps (199 yield) whereas the synthesis of their tert-Bu analogs failed.
The meso-isomer prevailed (dr = 90/10). Chiral thiazolium salts were
prepared in two steps from 2-tert-butylaniline. The enantiomerically
pure

pure
thiazolium salt I was obtained from α-bromomenthone and
2-text-butylaniline (27% overall yield). In contrast to the imidazolium
salts, the thiazolium salts proved to be subtable catalysts in the
benzoin
condensation of benzaldehyde and in the intramol. Stetter reaction of
2-OCHC6H4OCH2CH:CHCO2Me. The best results obtained with catalyst I (20
mol %) were 85% (R)-PhCOCHPHON (40% ee) and 75% Me (R)-4-oxochroman-3acetate. The stereogenic axis of I is not configurationally stable in
the

catalytically active carbene intermediate. The catalyst is recovered as

mixture of diastereomeric atropisomers in a ratio of 70:30 to 75:25.
REFERENCE COUNT: 68 THERE ARE 68 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RX(20) OF 30 COMPOSED OF RX(3), RX(4) RX(20) 2 G + 2 H ===> N

Searched by Jason M. Nolan

ANSWER 11 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

N: CM 1 YIELD 45%

RX (3) RCT G 49673-33-8

| STAGE(1)
RGT	K 7439-93-2 Li		
SOL	109-99-9 THF		
CON	SUBSTAGE(1)	room temperature	0 deg C

STAGE (2) RCT H 75-15-0 CON 20 hours, room temperature

Page 294

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L62 ANSWER 11 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

STAGE(3)

RGT L 7732-18-5 Water

PRO I 727417-84-7, J 727417-85-8

NTE ultrasound stage 1

RX(4) RCT I 727417-84-7

STAGE(1)

RGT O 7601-90-3 HC104, P 937-14-4 MCPBA

SOL 109-99-9 THF, 7732-18-5 Water

CON SUBSTAGE(1) room temperature -> -78 deg C

STAGE(2)

SOL 60-29-7 Et20

CON 2 hours, room temperature

PRO N 727417-87-0

NTE stereoselective, dr for meso:dl 9:1
```

RX(21) OF 30 COMPOSED OF RX(3), RX(5)RX(21) 2 G + 2 H ===> N

L62 ANSWER 11 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

RX(3) RCT G 49673-33-8

STAGE(1)

RCT K 7439-93-2 Li

SOL 109-99-9 THF

CON SUBSTAGE(1) room temperature

SUBSTAGE(2) room temperature -> 0 deg C

STAGE(2)

RCT H 75-15-0

CON 20 hours, room temperature

STAGE(3)

RCT L 7732-18-5 Water

PRO I 727417-84-7, J 727417-85-8

NTE ultrasound stage 1

RX(5) RCT J 727417-85-8

STAGE(1)

RX(5) RCT O 7601-90-3 HC104, P 937-14-4 MCPBA

STAGE(1)

RGT 0 7601-90-3 HCl04, P 937-14-4 MCPBA
SOL 109-99-9 THF, 7732-18-5 Water
CON SUBSTAGE(1) room temperature -> -78 deg C
STAGE(2)
SOL 60-29-7 Et2O
CON 2 hours, room temperature

PRO N 727417-87-0
NTE stereoselective, dr for meso:dl 9:1

L62 ANSWER 11 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

L62 ANSWER 12 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 141:106210 CASREACT

TITLE: Studies on high-temperature amination reactions of aromatic chlorides using discrete Palladium-N-Heterocyclic Catchene (NHC) complexes and in situ palladium/imidazolium salt protocols

AUTHOR(S): McCarroll, Andrew J.; Sandham, David A.; Titcomb,

Lisa R.; Lewis, Alexandra K. de K.; Cloke, F. Geoffrey N.;

Davies, Brian P.; Perez de Santana, Alejandro;

Hiller, Wolfgang; Caddick, Stephen

CORPORATE SOURCE: Physics and Environmental Sciences, School of Chemistry, Chemistry Laboratory, University of

Sussex,

Brighton, Falmer, UK

Molecular Diversity (2003), 7(2-4), 115-123

CODEN: MODIF4; ISSN: 1381-1991

PUBLISHER: Kluwer Academic Publishers

JOURNAL English

AB The palladium catalyzed coupling of aryl chlorides and amines can be readily achieved with short reaction times when carried out at high temps.

under thermal or microwave conditions. These coupling protocols are successful using two coordinate palladium-N-heterocyclic carbene complexes, or imidazolium salt protocols. Thus, Pd(dba)2/1,3-bis(disopropylphenyl)imidazolium tetrafluoroborate catalyzed coupling reaction of 4-HeC6MGCl with morpholine in the presence of KOBu-t in DME/OMF in microwave oven gave 97t N-(4-methylphenyl)morpholine.

REFERENCE COUNT: 62 THERE ARE 62 CITED REFERENCES AVAILABLE FOR THES

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L62 ANSWER 12 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

i-Pr

• c1-

C YIELD 35%

RX(1) RCT A 3188-13-4, B 74663-75-5 PRO C 250285-32-6 CAT 7732-10-5 Water SOL 109-99-9 THF CON 16 hours, room temperature

RX(17) OF 17 COMPOSED OF RX(1), RX(16) RX(17) A + B ===> B

L62 ANSWER 12 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

-F-B--F-

S: CM 1 YIELD 98

> Pr-i H i-Pr

S: CM 2 YIELD 98%

RX(1) RCT A 3188-13-4, B 74663-75-5 PRO C 250285-32-6 CAT 7732-18-5 Water SOL 109-99-9 THF CON 16 hours, room temperature

RX(16) RCT C 250285-32-6 RGT AM 13826-83-0 NH4.BF4 PRO S 282109-03-5 SOL 7732-18-5 Water CON room temperature

L62 ANSWER 13 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 140:217168 CASREACT
TITLE: Some Heterocyclization Reactions of
N,N'-Dimethoxycarbonyl-o-benzoquinone Diimine
AUTHOR(S): Velikorodov, A. V.; Babaitsev, D. D.; Mochalin, V. B.
CORPORATE SOURCE: Astrakhan State Pedagogical University, Astrakhan,
414056, Russia
SOURCE: Russian Journal of Organic Chemistry (Translation of
Enural Organicheskoi Khimii) (2003), 39(8),

1200-1201

PUBLISHER: MAIK Nauka/Interperiodica Publishing
DOCUMENT TYPE: Journal
LANGUAGE: English
AB Diels-Alder reaction with reversed electronic requirements were reported
on title compound reaction with RHC:CHR1 (R = C6H5; R1 = H; RR1 =
CH2CH2CH2CH2CH2).
REFERENCE COUNT: 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR
THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

...2.

RX(4) RCT I 2932-82-3, B 139499-83-5 PRO J 664333-94-2 SOL 67-66-3 CRC13, 60-29-7 Et2O CON 1 hour, 0 - 5 deg C L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

TITLE:

Aymmetric addition of aryl boron reagents to enones with rhodium dicyclophane imidazolium carbene catelysis

AUTHOR(S):

AUTHOR(S):

CORPORATE SOURCE:

CORPORATE SOURCE:

Department of Chemistry and Biochemistry, Brigham Young University, Provo, UT, 84602-5700, USA

Angewandte Chemie, International Edition (2003), 42(47), 5871-5874

CODEN: ACIEFS: ISSN: 1433-7851

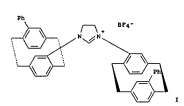
PUBLISHER:

DOCUMENT TYPE:

LANGUAGE:

English

GI



AB Chiral dicyclophane imidazoliumcarbene ligands, e.g., I, were prepared and screened as catalyst in addition reaction of cyclohexenone and arylboronic ends in the presence of rhodium. It was found that the ligand with 2-methoxyphenyl substituents on the dicyclophane gave the highest enantiomeric excess and isolation yield. The catalyst was used effectively in asym. conjugate addition of alkenones with arylboron reagents to yield chiral ketones in high yield.

REFERENCE COUNT: 51 THERE ARE 51 CITED REFERENCES AVAILABLE FOR THIS

RX(60) OF 91 COMPOSED OF RX(35), RX(36) RX(60) BE + N ===> BG L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

STEPS

ΒE

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RX(35) RCT BE 658711-12-7

STAGE(1)

RGT P 16940-66-2 NaBH4

SOL 109-99-9 THF

CON SUBSTAGE(1) 1 hour, 0 deg C

SUBSTAGE(1) 0 deg C -> room temperature

SUBSTAGE(3) 16 hours, room temperature

SUBSTAGE(4) 3 hours, reflux

SUBSTAGE(5) reflux -> room temperature

STAGE(2) RGT Q 7732-18-5 Water CON 0.5 hours

STAGE (3) RGT R 7647-01-0 HC1 PRO BF 658711-13-8

RCT BF 658711-13-8, N 122-51-0 PRO BG 658712-03-9 CAT 64-18-6 HCC2F CON SUBSTAGE(1) 60 hours, reflux SUBSTAGE(2) reflux -> room temperature RX (36)

RX(65) OF 91 COMPOSED OF RX(40), RX(43) RX(65) BH + N ===> EN

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

ві

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

● c1-

BO YIELD 90%

RX(41) RCT BI 658711-15-0

STAGE(1)

RGT P 16940-66-2 NaBH4

SOL 109-99-9 THF

CON SUBSTAGE(1) 1 hour, 0 deg C

SUBSTAGE(2) 0 deg C -> room temperature

SUBSTAGE(3) 16 hours, room temperature

SUBSTAGE(4) 3 hours, reflux

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RX (40)

RCT BH 658711-14-9 STAGE (1) AGE(1)
RGT P 16940-66-2 NABH4
SOL. 109-99-9 THF
CON SUBSTAGE(1) 1 hour, 0 deg C
SUBSTAGE(2) 0 deg C -> room temperature
SUBSTAGE(3) 16 hours, room temperature
SUBSTAGE(4) 3 hours, reflux
SUBSTAGE(4) 3 hours, reflux STAGE (2) RGT Q 7732-18-5 Water CON 0.5 hours STAGE(3) RGT R 7647-01-0 HC1 PRO BK 658711-17-2 RCT BK 658711-17-2, N 122-51-0
PRO BN 658712-06-2
CAT 64-18-6 HCOZH
CON SUBSTAGE(1) 60 hours, reflux
SUBSTAGE(2) reflux -> room temperature RX (43)

RX(66) OF 91 COMPOSED OF RX(41), RX(44) RX(66) BI + N ===> BO

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN SUBSTAGE(5) reflux -> room temperature (Continued)

> STAGE(2) RGT Q 7732-18-5 Water CON 0.5 hours STAGE (3) RGT R 7647-01-0 HC1 PRO BL 658711-18-3

BL 658711-18-3, N 122-51-0 BO 658712-09-5 64-18-6 HCO2H SUBSTAGE(1) 60 hours, reflux SUBSTAGE(2) reflux -> room temperature RX (44)

RX(67) OF 91 COMPOSED OF RX(42), RX(45) RX(67) BJ + N ===> BP

STEPS

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

06/28/2006 10/520,800

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

PAGE 2-A

● c1-

BP YIELD 90%

RX (42) RCT BJ 658711-16-1

> STAGE (1) RGT SOL CON

P 16940-66-2 NaBH4
109-99-9 THF
SUBSTAGE(1) 1 hour, 0 deg C
SUBSTAGE(2) 0 deg C -> room temperature
SUBSTAGE(3) 16 hours, room temperature
SUBSTAGE(4) 3 hours, reflux
SUBSTAGE(5) reflux -> room temperature

STAGE(2) RGT Q 7732-18-5 Water CON 0.5 hours

STAGE(3) RGT R 7647-01-0 HC1

PRO BM 658711-19-4

RCT BM 658711-19-4, N 122-51-0
PRO BP 658712-11-9
CAT 64-18-6 HCC2H
CON SUBSTAGE(1) 60 hours, reflux
SUBSTAGE(2) reflux -> room temperature RX (45)

RX(75) OF 91 COMPOSED OF RX(35), RX(36), RX(52) RX(75) BE + N ===> O

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN PRO BG 658712-03-9 CAT 64-18-6 HCO2H CON SUBSTAGE(1) 60 hours, reflux SUBSTAGE(2) reflux -> room temperature

RX (52)

BG 658712-03-9 S 13826-83-0 NH4.BF4 O 658711-04-7 67-56-1 HeOH SUBSTAGE(1) 3 hours, reflux SUBSTAGE(2) reflux -> room temperature

RX(83) OF 91 COMPOSED OF RX(40), RX(43), RX(46) RX(83) BH + N ===> W

STEPS

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

STEPS 0: CM 1 0: CM 2

RX (35) RCT BE 658711-12-7

STAGE(1)
RGT P 16940-66-2 NaBH4
SOL 109-99-9 THF
CON SUBSTAGE(1) 1 hour, 0 deg C
SUBSTAGE(2) 0 deg C -> room temperature
SUBSTAGE(2) 1 hours, room temperature
SUBSTAGE(4) 3 hours, reflux
SUBSTAGE(5) reflux -> room temperature

STAGE(2) RGT Q 7732-18-5 Water CON 0.5 hours

STAGE (3) RGT R 7647-01-0 HC1 PRO BF 658711-13-8

RX (36) RCT BF 658711-13-8, N 122-51-0

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

W: CM 2

RX (40) RCT BH 658711-14-9

STAGE(1)

RGT P 16940-66-2 NaBH4

SOL 109-99-9 THF

CON SUBSTAGE(1) 1 hour, 0 deg C

SUBSTAGE(2) 0 deg C -> room temperature

SUBSTAGE(2) 1 hours, room temperature

SUBSTAGE(3) 13 hours, reflux

SUBSTAGE(5) reflux -> room temperature

STAGE(2) RGT Q 7732-18-5 Water CON 0.5 hours

STAGE (3) RGT R 7647-01-0 HC1

PRO BK 658711-17-2

RCT BK 658711-17-2, N 122-51-0
PRO BN 658712-06-2
CAT 64-18-6 HCOZH
CON SUBSTAGE(1) 60 hours, reflux
SUBSTAGE(2) reflux -> room temperature RX (43)

RCT BN 658712-06-2 RGT 9 13826-83-0 NN4.BF4 PRO W 659711-06-9 SOL 67-56-1 MeOH CON SUBSTAGE(1) 3 hours, reflux SUBSTAGE(2) reflux -> room temperature RX (46)

RX(85) OF 91 COMPOSED OF RX(41), RX(44), RX(47) RX(85) BI + N ===> X

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

Et 
$$0$$
Et  $0$ 
E

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

PAGE 2-A

X: CM 2

RX (41) RCT BI 658711-15-0

> STAGE (1) )
> P 16940-66-2 NABH4
> 109-99-9 THF
> SUBSTAGE(1) 1 hour, 0 deg C
> SUBSTAGE(2) 0 deg C -> room temperature
> SUBSTAGE(3) 16 hours, room temperature
> SUBSTAGE(4) 3 hours, reflux
> SUBSTAGE(5) reflux -> room temperature

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

PAGE 2-A

Y: CM 2

RX (42) RCT BJ 658711-16-1

> AGE(1)
> RGT P 16940-66-2 NABH4
> SOL 109-99-9 THF
> CON SUBSTAGE(1) 1 hour, 0 deg C
> SUBSTAGE(2) 0 deg C -> room temperature
> SUBSTAGE(3) 16 hours, room temperature
> SUBSTAGE(3) 3 hours, reflux
> SUBSTAGE(4) 3 hours, reflux
> SUBSTAGE(5) reflux -> room temperature STAGE (2)

RGT Q 7732-18-5 Water CON 0.5 hours STAGE(3) RGT R 7647-01-0 HCl

PRO BM 658711-19-4

RX (45)

RCT BM 658711-19-4, N 122-51-0
PRO BP 658712-11-9
CAT 64-19-6 HOCZH
CON SUBSTAGE(1) 60 hours, reflux
SUBSTAGE(2) reflux -> room temperature

BP 658712-11-9 \$ 13826-83-0 NH4.BF4 7 65871-10-5 67-56-1 MeOH SUBSTAGE(1) 3 hours, reflux SUBSTAGE(2) reflux -> room temperature RCT RGT PRO RX (48)

L62 ANSWER 14 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) STAGE (2) RGT Q 7732-18-5 Water CON 0.5 hours

STAGE (3) RGT R 7647-01-0 HC1

PRO BL 658711-18-3 RX (44)

RCT BL 658711-18-3, N 122-51-0
PRO BO 658712-09-5
CAT 64-18-6 HCCZH
CON SUBSTAGE(1) 60 hours, reflux
SUBSTAGE(2) reflux -> room temperature

RCT BO 658712-09-5 RGT S 13826-83-0 NH4.BF4 PRO X 658711-08-1 SOL 67-56-1 MeOH RX (47) SUBSTAGE(1) 3 hours, reflux SUBSTAGE(2) reflux -> room temperature

RX(87) OF 91 COMPOSED OF RX(42), RX(45), RX(48) RX(87) BJ + N ===> X

Et 
$$0$$
 Et  $0$  E

L62 ANSWER 15 OF 35

ACCESSION NUMBER:

TITLE:

AN \*\*-heterocyclic carbene ligand with flexible steric bulk allows Suxuk cross-coupling of sterically hindered aryl effortices at room/temperature Altenboff, pfecon Goddard, Rickard; Lehmann, Christian (; Glorius, Frank

CORPORATE SOURCE:

SOURCE:

Angewandte Cheat, 1470, Germany
Angewandte Cheat, International Edition (2003), 42(31), 3809-3693

CODEN: ACIEFS: ISSN: 1433-7851

DOCUMENT TYPE:
LANGUAGE:

Brilish

CASREACT COPYRIGHT 2006 ACS on STN

139:27684 CASREACT
AN \*\*-heterocyclic carbene ligand with flexible steric bulk allows Excited at Toom/temperature
Alternational Chemany
Angewandte Chemany
Angew

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

A catalyst prepared from Pd(0Ac)2 and imidazolium salt I catalyzed the Suzuki cross-coupling of sterically hindered and unhindered, activated

unactivated, aryl chlorides and aryl boronic acids. Obtained were di-

and tri-ortho-substituted biphenyl compds.

REFERENCE COUNT: 34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

...L + M + H ===> N RX(4) OF 23

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L62 ANSWER 15 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                            (Continued)
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L62 ANSWER 15 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

RCT C 101725-44-4
RGT G 7719-09-7 SOC12
PGC F 66970-66-5
SOL 108-88-3 PhMe
CON SUBSTAGE(1) 1 hour, 60 deg C
SUBSTAGE(2) 3 hours, 90 deg C RX (2)

F 606970-66-5
I 1310-73-2 NAOH
H 606970-67-6
109-99-9 THF, 64-17-5 EtOH
SUBSTAGE(1) 20 minutes, room temperature
SUBSTAGE(2) room temperature -> 80 deg C
SUBSTAGE(3) 1.5 hours, 80 deg C RX (3)

RX (4) RCT L 2923-28-6, M 18997-19-8 STAGE(1)

SOL 75-09-2 CH2C12 CON 45 minutes STAGE (2) RCT H 606970-67-6 CON 20 hours, 40 deg C

PRO N 605970-69-8 NTE in the dark second stage

RCT F 606970-66-5
RGT I 1310-73-2 NAOH
PRO H 606970-671-6
SUL 109-99-9 THF, 64-17-5 EtOH
CON SUBSTAGE (1) 20 minutes, room temperature
SUBSTAGE (2) room temperature -> 80 deg C
SUBSTAGE (3) 1.5 hours, 80 deg C RX (3)

RX (4) RCT L 2923-28-6, M 18997-19-8 STAGE(1) SOL 75-09-2 CH2C12 CON 45 minutes

STAGE(2) RCT H 606970-67-6 CON 20 hours, 40 deg C N 606970-69-8 in the dark second stage

RX(22) OF 23 COMPOSED OF RX(2), RX(3), RX(4) RX(22) C + L + M ===> N

L62 ANSWER 16 OF 35 CASREACT COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 139:276460 CASREACT TITLE: Sonogashira Coupling Using Bulky Palladium-Phenanthryl

Palladium-Phenanthryl

AUTHOR(S):

Ma, Yudao; Song, Chun; Jiang, Wei; Wu, Quansheng; Wang, Yong; Liu, Xueying; Andrus, Herritt B.

CORPORATE SOURCE:

Department of Chemistry and Biochemistry, Brigham Young University, Provo, UT, 84602-5700, USA

Organic Letters (2003), 5(18), 3317-3319

COUNTRY TYPE:

DOCUMENT TYPE:

LANGUAGE:

BUlky phenanthrenyl imidazolium-derived carbene ligands were investigated for copper-free Sonogashira coupling with terminal acetylenes. Aryl bromides and iodides gave coupled products in excellent yields from the Pd(PPh3)2C12 complex with potassium t-butoxide and 18-crown-6 in THF. A remarkable dependence on the size of the ligand was found. The highest yields were obtained with the bulky (2,9-dicyclohexyl-10-phenanthrenyl)imidazolylidene ligand.

REFERENCE COUNT:

32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THERE

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

RX(53) OF 74 COMPOSED OF RX(44), RX(37) RX(53) CM + BG sum> G

K

L62 ANSWER 16 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

STEPS

. STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT .

RCT CE 605686-25-7
RGT CH 16940-66-2 NaBH4
PRO BF 605686-26-8
SOL 109-39-9 THF
CON SUBSTAGE(1) 0 deg C
SUBSTAGE(2) 1 hour, 0 deg C
SUBSTAGE(3) 0 deg C -> room temperature
SUBSTAGE(4) 16 hours, room temperature
SUBSTAGE(5) 3 hours, reflux RX (44)

BF 605686-26-8, BG 122-51-0 G 605686-20-2 64-18-6 HCO2H 60 hours, reflux RX (37)

L62 ANSWER 17 OF 35
ACCESSION NUMBER:
139:6825 CASREACT
TITLE:
New N-acyl, N-alkyl, and N-bridged derivatives of rac-6,6',7,'-tetramethoxy-1,1',2,2',3,3',4,4'-octahydro-1,1'-brisisoquinoline
AUTHOR(S):
Busato, Stephan; Craig, Donald C.; Judeh, Zaher M.

AUTHOR(S): A.;

Acir Susaco, Stephanic Craig, bonaid C., duden, Zaner H.

CORPORATE SOURCE: School of Chemical Sciences, The University of New South Wales, Sydney, 2052, Australia Tetrahedron (2003), 59(4), 461-472

CODEN: TETRAB: ISSN: 0040-4020

Elsevier Science Ltd.

DOCUMENT TYPE: English

AB The preparation of potential new ligand systems based on the rac-1,1',2,2',3,3',4,4'-octahydro-6,6',7,7'-tetramethoxy-1,1'-bisioquinoline skeleton has been investigated. Syntheses of N-(2-bromobensyl), N-(3-acetoxybensyl), N-acetyl, N-chlorocactyl, N-chlorocarbonyl, N-chlorocarbonyl and N-tetr-butyloxycarbonyl derivs.

and
five macrocyclic, polyether containing derivs. are also described. Asym.
reduction of one of the bisamine compound is also reported. Crystal
structure
of some of the products were also investigated.

REFERENCE COUNT: 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR
THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

RX(44) OF 52 COMPOSED OF RX(14), RX(15) RX(44) 2 AO + M ===> AT

L62 ANSWER 17 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

AT YIELD 94%

RX (14)

STAGE(1)
RGT AQ 1148-11-4 1,2-Pyrrolidinedicarboxylic acid,
1-(phenylmethyl) ester, (2S)-, AR 25895-60-7 NaBH3CN
SOL 109-99-9 THF
CON 0 deg C STAGE(2)
RCT AO 30340-61-5
SOL 109-99-9 THF
CON SUBSTAGE(1) -25 deg C
SUBSTAGE(2) 15 hours, -25 deg C -> room temperature

RCT M 75-09-2, B 75370-82-0 RGT P 584-08-7 K2CO3 PRO AT 75370-84-2 CON 3 days, reflux RX (15)

ACCESSION NUMBER:

138:321365 CASREACT

TITLE:

OARZOÎINES as chiral building blocks for imidazolium salts and N-heterocyclic carbene ligands

Glorius, Frank: Altenhoff, Gereon; Goddard, Richard;

Lehmann, Christian

ORPORATE SOURCE:

MAX-Planck-Institut fuer Kohlenforschung,
Muelheim/Ruhr, 45470, Germany

COBEN:

CODEN: CHCOPS; ISSN: 1359-7345

ROUBLISHER:

ROYAL SOCIETY Of Chemistry

DOCUMENT TYPE:

JOURNAL SOCIETY OF Chemistry

DOCUMENT TYPE:

JOURNAL SOCIETY OF Chemistry

DOCUMENT TYPE:

AB Enantiomerically pure imidazolium triflates can be readily prepared from bioxazolines and oxazolinemines. Deprotonation of imidazolium triflate gives a chiral N-heterocyclic carbene that can act as a ligand in a catalytically active palladium complex.

REFERENCE COUNT:

22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

RX(1) OF 13 A + B ===> C...

C: CM 1 YIELD 80% C: CM 2 YIELD 80%

RCT A 131833-89-1, B 18997-19-8 RGT D 2923-28-6 AgO3SCF3 PRO C 512193-98-5 SOL 75-09-2 CH2C12 CON 24 hours, 40 deg C NTE in the dark RX (1)

F + B ===> G RX(2) OF 13

×

L62 ANSWER 18 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

G: CM 1 YIELD 75% G: CM 2 YIELD 75%

F 135565-31-0, B 18997-19-8 D 2923-28-6 AgO3SCF3 G 512194-01-3 75-09-2 CH2Cl2 24 hours, 40 deg C in the dark RCT RGT PRO RX (2)

RX (3) OF 13 B + H ===> I

RCT B 18997-19-8 RX (3)

L62 ANSWER 18 OF 35 CASREACT COPYRIGHT 2006 ACS on STN STAGE(1) (Continued) D 2923-28-6 AgO3SCF3 75-09-2 CH2C12 1 hour RGT SOL CON

STAGE (2) RCT H 133463-88-4 CON 24 hours, 40 deg C

PRO I 512194-04-6 NTE in the dark

RX(12) OF 13 COMPOSED OF RX(1), RX(8) RX(12) A + B ===> AC

AC

A 131833-89-1, B 18997-19-8 D 2923-28-6 AgO3SCF3 C 512193-98-5 75-09-2 CH2C12 24 hours, 40 deg C in the dark RCT RGT PRO RX (1) SOL

C 512193-98-5 AD 7693-26-7 KH AC 512194-17-1 865-47-4 t-BuoK 109-99-9 THF RCT RGT PRO RX (8)

L62 ANSWER 19 OF 35
ACCESSION NUMBER:
TITLE:
Optically Active Iridium Imidazol-2-ylidene-oxazoline
Complexes: Preparation and Use in Asymmetric
Hydrogenation of Arylalkenes
Perry, Marc C.; Cui, Xiuhua; Powell, Mark T.; Hou,
Duen-Ren; Reibenspies, Joseph H.; Burgess, Kevin
Chemistry Department, Texas A H University, College
Station, TX, 77842, USA
JOURNAL OF THE American Chemical Society (2003),
125(11, 113-123
CODEN: JACSART; ISSN: 0002-7863
American Chemical Society
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
English

PUBLISHER DOCUMENT LANGUAGE: GI

т

A library of iridium imidazolylidene oxazoline complexes [I: wherein M = Ir: Rl = 1-Ad, t-Bu, CHPh2, Ph, etc.; R2 = t-Bu, CHPh2, Cy,

AB A library of iridium imidazolylidene oxazoline complexes [I, wherein M = Ir; R1 = 1-Ad, t-Bu, CHPh2, Ph, etc.; R2 = t-Bu, CHPh2, Cy,
2.4.6-Me3C6H2,
3.5-t-Bu2-4-MeOC6H2, 2.5-Et2C6H3, 2.6-i-Pr2-C6H3, 2.5-t-Bu2-C6H3, 1-Ad, etc.] were prepared and used as catalysts in asym. hydrogenations of arylalkenes. Three of the complexes [M = Ir; R1 = 1-Ad, R2 = t-Bu (Sab); R1 = t-Bu (Sab);

L62 ANSWER 19 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) enantiomeric excess ys. +89% enantiomeric excess); a transformation from one prevalent mechanism to another is inferred from this. The studies of pressure dependence revealed that many reactions proceeded with high conversions, and optimal enantioselectivities in approx. 2 h when only 1 bar of hydrogen was used. Deuterium-labeling expts. provide evidence for other types of competing mechanisms that lead to D-incorporation at positions that do not correspond to direct addn. to the double bond.

REFERENCE COUNT: 74 THERE ARE 74 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

RX(67) OF 143 COMPOSED OF RX(14), RX(22) RX(67) AP + AF + BH ===> BL

STEPS

BL

RCT AP 343217-35-6, AF 288-32-4 RX (14)

AH 534-17-8 Cs2CO3, AI 538-58-9 1,4-Pentadien-3-one, 1,5-diphenyl-, AJ 66-71-7 1,10-Phenanthroline, AK

X

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L62 ANSWER 19 OF 35 CASREACT COPYRIGHT 2006 ACS on STN 42152-44-3 Cuprous triflate
SOL 1330-20-7 Kylene
CON SUBSTAGE[1] 36 hours, 125 deg C
SUBSTAGE[2] 125 deg C -> room temperature
                                                                                                                  (Continued)
                      STAGE (2)
                            RGT F 12125-02-9 NH4C1
SOL 75-09-2 CH2C12, 7732-18-5 Water
                   PRO AO 496067-55-1
                  RCT AQ 496067-55-1, BH 369657-19-2
PRO BL 496067-64-2
SOL 68-12-2 DMF
CON 12 hours, 80 deg C
RX (22)
                           stereoselective
RX(108) OF 143 COMPOSED OF REACTION SEQUENCE RX(28), RX(22) AND REACTION SEQUENCE RX(14), RX(22)
            ===> BH...
+ AF + BH ===> BL
                                                                                    STEPS
```

START NEXT REACTION SEQUENCE

L62 ANSWER 19 OF 35 CASREACT COPYRIGHT 2006 ACS on STN STAGE(2)

RCT F 12125-02-9 NH4C1
SOL 75-09-2 CH2C12, 7732-18-5 Water PRO AQ 496067-55-1 RCT AQ 496067-55-1, BH 369657-19-2 PRO BL 496067-64-2 SOL 68-12-2 DMF CON 12 hours, 80 deg C NTE stereoselective RX (22)

L62 ANSWER 19 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) AP STEPS ● T · ВL RCT BV 369657-20-5 RGT BW 7681-11-0 KI PRO BH 369657-19-2 SOL 67-64-1 Me2CO RX (28) 4 hours, 55 deg C RX (14) RCT AP 343217-35-6, AF 288-32-4 )
AM 534-17-8 Cs2CO3, AI 538-58-9 1,4-Pentadien-3-one,
1,5-diphenyl-, AJ 66-71-7 1,10-Phenanthroline, AK
42152-44-3 Cuprous triflate
1330-20-7 Xylene
SUBSTAGE(1) 36 hours, 125 deg C
SUBSTAGE(2) 125 deg C -> room temperature

L62 ANSWER 20 OF 35
ACCESSION NUMBER:
TITLE:
Synthesis of a transient tropylidene substituted
N-heterocyclic carbene (tropNHC): rearrangement and
formation of its gold complex
Boehler, Carsten: Stein, Daniel: Donati, Nicola;
Gruetzmacher, Hansjoerg
CORPORATE SOURCE:
Department of Chemistry, Laboratory of Inorganic
Chemistry, ETH-Neonggerberg, Zurich, CH-8093, Switz.
New Journal of Chemistry (2002), 26(10), 1291-1295
CODEN: NJCHES: ISSN: 1144-0546
PUBLISHER:
ROYAL Society of Chemistry
LANCUAGE:
English
BT the condensation reaction of the primary tropylidenyl amine tropamine
RNH2 AB The Condensation reaction of the primary tropylidenyl amine tropamine RNNI2

(2, R = 5H-dibenzo[a,d]cyclohepten-5-yl) with glyoxal 3 leads to the corresponding 1,4-diazadiene bistropdad RN:CKCH:NR (4) in high yield. With formaldehyde and ethereal HCl, 4 is transformed to the bistropimidazolium salt 1,3-R2-imidazolium chloride (5). Deprotonation with KOLBU in THF did not gave a stable N-heterocyclic carbene bistropNHC 1,3-R2-imidazol-2-ylidene (6), but the imidazole derivative 2-(5H-dibenzo[a,d]cyclohepten-10-yl)-1-R-IN-imidazole 9 as a product of a rearrangement. However, the unstable carbene 6 can be trapped when it is generated in the presence of [AuCl(PPh3)] whereby the stable cationic mixed phosphine carbene gold complex (1,3-R2-imidazol-2-ylidene)[PPh3]aulCl (10) was obtained and characterized by x-ray diffraction.

REFERRNCE COUNT: 31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

RX(2) OF 9 ...C + F ===> G...

FORMAT

н₂с≐о (2)

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L62 ANSWER 20 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                           (Continued)
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G YIELD 79%

RCT C 492446-84-1, F 50-00-0 RX (2)

STAGE(1) SOL 108-88-3 PhMe CON room temperature

STAGE(2)
RGT H 7647-01-0 HC1
SOL 60-29-7 Et20
CON 2 days, room temperature

PRO G 492446-85-2 NTE paraformaldehyde was used

L62 ANSWER 21 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 138:136761 CASREACT
TITLE: Acylotropic Tautomerism: XXXV. R.dblarw.L-Inversion
of

Configuration of Dipolar Spyrocyclic and Open-Chain 2-Arylaminotropone Isomers
Olekhnovich, L. P.; Budarina, Z. N.; Borodkin, G. S.;
Kurbatov, S. V.; Vaslyaeva, G. S.; Zhdanov, Yu. A.
Rostov State University, Rostov-on-Don, 344090, AUTHOR (S):

NOSTOV State University, Rostov-on-Don, 344090,

NUSSIA

SOURCE: Russian Journal of Organic Chemistry (Translation of Zhurnal Organicheskoi Khimii) (2002), 38(5), 713-722 CODEN: RJOCEQ; ISSN: 1070-4280

PUBLISHER: MAIK Nauks/Interperiodica Publishing Journal LANGUAGE: Journal LANGUAGE: Journal LANGUAGE: A Ribiary L-Inversion of chiral spirocyclic and open-chain 2-arylaminotropone derivs. With varied heteroatom (0, S, N) was studied. Kinetic relations holding in the RI-permutation are discussed. Its mechanism includes formation and dissociation of spiro bonds and torsion stereodynamics.

REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILable This

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

VERIFICATION INCOMPLETE

RX(8) OF 35 ...S + B ===> T

L62 ANSWER 21 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

T YIELD 66%

RX (8)

RCT S 491879-92-6, B 954-50-7
PRO T 492435-83-3
SOL 67-66-3 CRCl3
CON SUBSTAGE (1) room temperature -> reflux
SUBSTAGE (2) reflux -> 80 deg C
SUBSTAGE (3) 6 - 8 hours, 80 deg C

VERIFICATION INCOMPLETE

RX(9) OF 35 ...U + B ===> V

L62 ANSWER 21 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

V YIELD 78%

RX (9)

RCT U 491879-91-5, B 954-50-7
PRO V 492435-84-4
SOL 67-66-3 CRC13
CON SUBSTAGE (1) room temperature -> reflux
SUBSTAGE (2) reflux -> 80 deg C
SUBSTAGE (3) 6 - 8 hours, 80 deg C

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 137:353265 CASREACT
TITLE: Convenient synthesis of human calcitonin and its methionine sulfoxide derivative
AUTHOR(S): Shi, Tiesheng: Rabenstein, Dallas L.
CORPORATE SOURCE: Department of Chemistry, University of California, Riversade, CA, 92521, USA
SOURCE: Bioorganic & Medicinal Chemistry Letters (2002), 12(16), 2237-2240
CODEM: EMCLE8: ISSN: 0960-894X
FUBLISHER: Elsevier Science Ltd.
DOCUMENT TYPE: Journal
LANGUAGE: English
AB The human calcitonin peptide chain was assembled using Pmoc solid-phase peptide synthesis chemical The combinations of cleavage Reagent H (TFA 811,

phenol 5%, thioanisole 5%, ethanedithiol 2.5%, dimathylsulfide 2%, water 3%, ammonium iodide 1.5%) with trans-{Pt(en)2Cl2|2+ and Reagents B (TFA 88%, phenol 5%, triisopropylsilane 2%, and water 5%), K (TFA 82.5%, phenol 5%, thioanisole 5%, ethanedithiol 2.5%, and water 5%), and R (TFA 90%, thioanisole 5%, ethanedithiol 3%, anisole 2%) with trans-[Pt(CN)4Cl2]2-provide convenient methods for the synthesis of human calcitonin and its methionine sulfoxide derivative; the formation of intramol. disulfide s by

methionine sulfoxide derivative; the formation of intramol. disulfide bonds by the above Pt(IV) oxidants is essentially quant.

REFERENCE COUNT: 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

VERIFICATION INCOMPLETE

RX(40) OF 40 COMPOSED OF REACTION SEQUENCE RX(1), RX(3), RX(6)
AND REACTION SEQUENCE RX(2), RX(3), RX(6)
AND REACTION SEQUENCE RX(1), RX(4), RX(6)
AND REACTION SEQUENCE RX(1), RX(4), RX(6)
...A + B + C + D + E + F + G + H + I + J + N + O + P + Q + X ===> Y... B + C + D + E + F + G + H + I + 

STEPS

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

PAGE 2-D

START NEXT REACTION SEQUENCE

CO2H STEPS

START NEXT REACTION SEQUENCE

STRUCTURE STRUCTURE DIAGRAM DIAGRAM IS NOT IS NOT AVAI LABLE AVAI LABLE STEPS

START NEXT REACTION SEQUENCE

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

PAGE 1-A

- \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT \*
- \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT \*

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 AGS on STN (Continued)
STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

PAGE 2-D

3 STEPS

PAGE 1-A

10/520,800 06/28/2006

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

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-N
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RX(1) RCT A 71989-31-6

STAGE(1)

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-Prn:C:NPr-i

SOL 68-12-2 DMF

(Continued)

STAGE(2) RGT U 110-89-4 Piperidine SOL 68-12-2 DMF

STAGE(3)
RCT B 35661-39-3
RGT S 39968-33-7 3H-1,2,3-Triazolo{4,5-b}pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF

STAGE(4)
RCT C 29022-11-5
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF

STAGE(5)
RCT D 68858-20-8
RGT S 39968-33-7 3H-1,2,3-Triazolo{4,5-b}pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF

STAGE(6) RCT E 71989-23-6 RGT S 39968-33-7 3H-1,2,3-Triazolo(4,5-b]pyridine, 3-hydroxy-, T 693-13-0 i-PrN:C:NPr-i SOL 68-12-2 DMF

STAGE(7)
RCT F 73731-37-0
RGT S 39968-33-7 3H-1,2,3-Triazolo(4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) STAGE (8) RCT G 132327-80-1 RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-, T 693-13-0 i-PrN:C:NPr-i SOL 68-12-2 DMF STAGE (9) RGT U 110-89-4 Piperidine STAGE (10) RCT H 35661-40-6 RCT H 35661-40-6 RCT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-, T 693-13-0 i-PrN:C:NPr-i SOL 68-12-2 DMF STAGE(11) RGT U 110-89-4 Piperidine STAGE (12) RCT I 109425-51-6 RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-, T 693-13-0 i-PrN:C:NPr-i SOL 68-12-2 DMF STAGE (13)

RCT J 105047-45-8

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMF STAGE (15) RCT L 71989-14-5 RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-, T 693-13-0 i-PrN:C:NPr-i SOL 68-12-2 DMF STAGE (16) NOC.10)
RCT M 71989-38-3
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 66-12-2 DMF STAGE(17)
RCT N 35661-60-0
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF

L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN SOL 68-12-2 DMF (Continued) STAGE(19)
RCT P 103213-32-7
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF STAGE (20) RGT Q 71989-33-8 RGT Q 71989-33-7 RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-, T 693-13-0 i-PrN:C:NPr-i SOL 68-12-2 DMF PRO R 474527-92-9D
NTE solid-supported reaction, first stage is deprotection of FmocPAL-PEG-PS resin, std. side chains protecting groups
(tBu,trityl,Boc) assumed, piperidine used for all subsequent
deprotection after coupling R 474527-92-9D, X 474527-93-0D Z 76-05-1 F3CCO2H, AA 108-95-2 PhOH, AB 100-68-5 PhSMe, AC \$40-63-6 HSCHZCHZSH Y 27686-18-6 76-05-1 F3CCO2H solid-supported reaction, other products also detected RX (3) RX (2) RCT A 71989-31-6 STAGE(2) RGT U 110-89-4 Piperidine SOL 68-12-2 DMF STAGE (3) RCT B 35661-39-3 RCT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-, T 693-13-0 i-PrN:C:NPr-i SOL 68-12-2 DMF STAGE (4) AGE(4) RCT C 29022-11-5 RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-, T 693-13-0 i-PrN:C:NPr-i SOL 68-12-2 DMF STAGE(5) RCT D 68858-20-8 RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-, T 693-13-0 i-PrN:C:NPr-i SOL 68-12-2 DMF RCT E 71989-23-6 RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

06/28/2006

```
L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued) T 693-13-0 i-PN:C:NPr-i SOL 68-12-2 DMF
                                                                                                                                                                  L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN SOL 68-12-2 DMF
                                                                                                                                                                                                                                                                                (Continued)
                                                                                                                                                                                        STAGE(16)
RCT W 76265-70-8
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                      STAGE (7)
                           NOST 17 73731-37-0

RGT F 73731-37-0

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMF
                                                                                                                                                                                         STAGE (17)
                                                                                                                                                                                              NOSI17)
RCT P 103213-32-7
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                      STAGE (8)
                           AGE(8)
RCT G 132327-80-1
RGT S 39968-33-7 3H-1,2,3-Triazolo(4,5-b)pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                                                                                                                                                                                         STAGE (18)
                     RCT Q 71989-33-8
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                                                                                                                                                                                    PRO X 474527-93-0D
NTE solid-supported reaction, first stage is deprotection of Fmoc-
PAL-PEG-PS resin, std. side chains protecting groups
(tBu,trity),Boc) assumed, piperidine used for all subsequent
deprotection after coupling
                     STAGE(10)
RCT I 109425-51-6
RCT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 1-Prn:C:NPr-i
SOL 68-12-2 DMF
                                                                                                                                                                  RX (2)
                                                                                                                                                                                    RCT A 71989-31-6
                     STAGE(11)

RCT J 105047-45-8

RCT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-prn:C:NPr-i

SOL 68-12-2 DMF
                                                                                                                                                                                        STAGE (1)
                                                                                                                                                                                             RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                     STAGE(12)

RCT K 132388-59-1

RCT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMF
                                                                                                                                                                                        STAGE(2)
RGT U 110-89-4 Piperidine
SOL 68-12-2 DMF
                                                                                                                                                                                        STAGE(3)
RCT B 35661-39-3
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF.
                     STAGE(13)
RCT L 71989-14-5
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 1-PrN:C:NPr-i
SOL 66-12-2 DMF
                                                                                                                                                                                        STAGE(4)
RCT C 29022-11-5
RGT S 39968-33-7 3H-1,2,3-Triazolo(4,5-b)pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 66-12-2 DMF
                      STAGE(14)

RCT M 71989-38-3

RCT S 39966-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 1-PrN:C:NPr-i

SOL 68-12-2 DMF
                                                                                                                                                                                        STAGE(5)
RCT D 60858-20-8
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                     STAGE(15)
RCT N 35661-60-0
RGT S 39568-33-7 3H-1,2,3-Triazolo(4,5-b)pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                                                                                  L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                              (Continued)
                           GGE(6)
RCT E 71989-23-6
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 1-PrN:C:NPr-i
SOL 66-12-2 DMF
                                                                                                                                                                                        STAGE(16)
RCT W 76265-70-8
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                      STAGE (7)
                                                                                                                                                                                        STAGE(17)
RCT P 103213-32-7
ROT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                           RCT F 73731-37-0
RCT S 39568-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                      STAGE (8)
                                                                                                                                                                                        STAGE(18)

RCT Q 71989-33-8

RCT S 39968-33-7 3H-1,2,3-Triazolo(4,5-b)pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMF
                           RCT G 132327-80-1
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 60-12-2 DMF
                      STAGE (9)
                                                                                                                                                                                     PRO X 474527-93-0D
NTE solid-supported
                           RCT H 35661-40-6
RCT S 39568-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PFN:C:NPr-i
SOL 68-12-2 DMF
                                                                                                                                                                                              X 474527-93-0D solid-supported reaction, first stage is deprotection of Fmoc-
PAL-PEG-PS resin, std. side chains protecting groups (tBu,trityl,Boc) assumed, piperidine used for all subsequent deprotection after coupling
                      STAGE (10)
                           RCT I 109425-51-6
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                                                                                                                                                                  RX (1)
                                                                                                                                                                                         STAGE (11)
                           MCT J 105047-45-8

RGT J 105047-45-8

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 1-PrN:C:NPr-i

SOL 68-12-2 DMF
                                                                                                                                                                                         STAGE(2)
RGT U 110-89-4 Piperidine
SOL 68-12-2 DMF
                                                                                                                                                                                        STAGE(3)

RCT B 35661-39-3

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMF
                      STAGE (12)
                           NOCITY

RCT K 132388-59-1

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMP
                                                                                                                                                                                        STAGE(4)

RCT C 29022-11-5

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMF
                      STAGE (13)
                           AGE[13]

RCT L 71989-14-5

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 60-12-2 DMF
                                                                                                                                                                                         STAGE(5)

RCT D 68858-20-8

RCT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-prN:C:NPr-i

SOL 68-12-2 DMF
                     STAGE(14)

RCT M 71989-38-3

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMF
                                                                                                                                                                                         STAGE(6)
RCT E 71989-23-6
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                      STAGE(15)
RCT N 35661-60-0
RGT S 39568-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 1-PrN:C:NPr-i
SOL 68-12-2 DMF
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L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)
STAGE(17)
RCT N 35661-60-0
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PN:C:NPr-i
SOL 68-12-2 DMF
L62 ANSWER 22 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
                                                                                                                         (Continued)
                       STAGE (7)
                            RCT F 73731-37-0
RCT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                                                                                                                                                                                                      STAGE(18)

RCT 0 71989-28-1

RCT 5 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMF
                       STAGE (B)
                            AGE(8)
RCT G 132327-80-1
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                                                                                                                                                                                                      STAGE(19)
RCT P 103213-32-7
RCT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                      STAGE(9)
RGT U 110-89-4 Piperidine
                      STAGE(10)

RCT H 35661-40-6

RCT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMF
                                                                                                                                                                                                      STAGE(20)
RCT Q 71989-33-8
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                      STAGE(11)
RGT U 110-89-4 Piperidine
                                                                                                                                                                                                           R 474527-92-9D solid-supported reaction, first stage is deprotection of Fmoc-PAL-PEG-PS resin, std. side chains protecting groups (tBu, trityl,Boc) assumed, piperidine used for all subsequent deprotection after coupling
                       STAGE (12)
                            AGE(12)
RCT I 109425-51-6
RGT S 39968-33-7 3H-1,2,3-Triazolo(4,5-b)pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                                                                                                                                                                               RX (4)
                                                                                                                                                                                                   RCT R 474527-92-9D, X 474527-93-0D
RGT Z 76-05-1 F3CCO2H, AB 100-68-5 PhSMe, AC 540-63-6 HSCH2CH2SH,
                       STAGE (13)
                            AGE[13]
RCT J 105047-45-8
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 60-12-2 DMF
                                                                                                                                                                                                             100-66-3 PhoMe
AD 73840-80-9
76-05-1 F3CCO2H
solid-supported reaction, other products also detected
                       STAGE (14)
                                                                                                                                                                                                            Y 27686-18-6, AD 73840-80-9
AI 12072-77-4 Platinate(2-), dichlorotetrakis(cyano-KC)-,
dipotassium, (0C-6-12)-
AH 67881-33-8
buffered soln.
                            AGE[14]
RCT K 132388-59-1
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                                                                                                                                                                               RX (6)
                       STAGE (15)
                            WORLIS]
RCT L 71989-14-5
RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,
T 693-13-0 i-PrN:C:NPr-i
SOL 68-12-2 DMF
                       STAGE (16)
                            NGCT M 71989-38-3

RGT M 71989-38-3

RGT S 39968-33-7 3H-1,2,3-Triazolo[4,5-b]pyridine, 3-hydroxy-,

T 693-13-0 i-PrN:C:NPr-i

SOL 68-12-2 DMP
```

L62 ANSWER 23 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
137:140582 CASREACT
Synthesis and Structural Features of Arduengo Carbene Complexes of Group 4 Metallocene Cations
AUTHOR(S):
Schwab,
Pia: Froehlich, Roland: Blacque, Olivier; Berke,
Heinz
CORPORATE SOURCE:
Organisch-Chemisches Institut, Universitaet Muenster,
Muenster, D-48149, Germany
OURCE:
Organisch-Chemisches Institut, Universitaet Muenster,
Muenster, D-48149, Germany
OURCE:
Organisch-Chemisches Institut, Universitaet Muenster,
Muenster, D-48149, Germany
OURCE:
OURC

solution, featuring symmetry-equivalent Cp rings and a pair of diastereotopic iso-Pr substituents as well as chemical differentiated imidazol-2-ylidene

substituents as well as chemical differentiated imidazol-2-ylidene C4H:C5H groups. The reaction of the ion pair [(Cp22rCH3)+(CH3B(C6F5)3-)] (7)

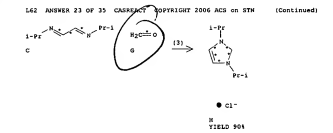
with 4 gave the analogous Arduengo carbene zirconocene cation complex (Cp2ZrMeL)MeB(C6F5)3 (6b; >95% isolated).

REFERENCE COUNT: 52 THERE ARE 52 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

RX(3) OF 23 ...C + G ===> H...

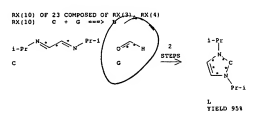


RX(3) RCT C 24764-90-7, G 50-00-0

STAGE (1)
SOL 108-88-3 PhMe

STAGE (2)
RGT I 7647-01-0 HC1
SOL 123-91-1 Dioxane

PRO H 139143-09-2 NTE paraformaldehyde was used



RX(3) RCT C 24764-90-7, G 50-00-0

STAGE(1)
SOL 108-88-3 PhMe

STAGE(2)
RGT I 7647-01-0 HC1
SOL 123-91-1 Dioxene

PRO H 139143-09-2

L62 ANSWER 23 OF 35 CASREACT COPYRIGHT 2006 ACS on STN NTE paraformaldehyde was used (Continued)

RX (4)

RCT H 139143-09-2 RGT M 7646-69-7 NaH, N 865-47-4 t-BUOK PRO L 179863-09-3 SOL 109-99-9 THF

L62 ANSWER 24 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
136:69622 CASREACT
Amination Reactions of Aryl Halides with
Nitrogen-Containing Reagents Mediated by
Palladium/Imidazolium Salt Systems
Grass, Gabriela A.; Viciu, Mihai S.; Huang, Jinkun;
Nolan, Steven P.
CORPORATE SOURCE:
Department of Chemistry, University of New Orleans,
New Orleans, LA, 70148, USA
Journal of Organic Chemistry (2001), 66(23),

SOURCE: 7729-7737

7729-737

CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER: American Chemical Society

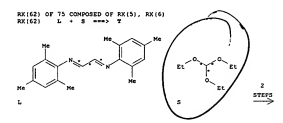
DOCUMENT TYPE: Journal

LANGUAGE: English

AB Nucleophilic N-heterocyclic carbenes have been conveniently used as catalyst modifiers in amination reactions involving aryl chlorides, aryl bromides, and aryl iodides with various nitrogen-containing substrates.

bromides, and aryl iodides with various nitrogen-containing substrates. The scope of a coupling process using a Pd(0) or Pd(II) source and an imidazolium salt in the presence of a base, KoCMe3 or NaOH, was tested using various substrates. The Pd2(dba)3/FP-HCI [IPF = 1,3-bis(2,6-diisopropylphenyl);midazol-2-ylidene) system presents the highest activity with respect to electron-neutral and electron-rich aryl chlorides. The ligand is also effective for the synthesis of benzophenone imines, which can be easily converted to the corresponding primary amines by acid hydrolysis. Less reactive indoles were converted to N-aryl-aubstituted indoles using as supporting ligand the more donating SIPr-HCI [SIPr = 1,3-bis(2,6-diisopropylphenyl)-4,5-dihydroimidazol-2-ylidenel. The Pd(Obal2/SIPr-HCI/MAOH system is efficient for the N-arylation of diverse indoles with aryl bromides. The general protocol developed has been applied successfully to the synthesis of a key intermediate in the synthesis of an important new antibiotic. Mechanistically, palladium-to-ligand ratio studies strongly support an active species bearing one nucleophilic carbene ligand.

THERE ARE 114 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT



L62 ANSWER 24 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RX (5)

RCT L 56222-36-7 RGT Q 16940-66-2 NaBH4 PRO P 134030-21-0 SOL 67-56-1 MeOH, 109-99-9 THF

RCT P 134030-21-0, S 122-51-0 RGT U 12125-02-9 NH4C1 PRO T 141556-45-8 RX (6)

L62 ANSWER 25 OF 35
ACCESSION NOMBER:
133:362823 CASREACT
133:3628

CODEN: JORCAI; ISSN: 0022-328X

PUBLISHER: Elsevier Science S.A.

DOCUMENT TYPE: Journal

LANGUAGE: English

B The sterically demanding nucleophilic carbene ligand 1,3-bis(2,6-disopropylphenyl)lmidazol-2-ylidene (IPr. 4) has been synthesized. The reaction of (Cp-RuCl]4 (S; Cp = n5-CMs6) with this ligand affords a coordinatively unsatd. Cp\*Ru(IPr)Cl (6) complex. Solution calorimetric results in this system provide information concerning the electron donor properties of the carbene ligand. Steric parameters associated with this ligand are determined from the x-ray crystal structure study. The carbene

ligand are determined from the x-ray crystal structure study. The carbene ligand reacts with RuCl2(:C(H)Ph)(PCy3)2 to yield a mixed carbene-phosphine ruthenium complex RuCl2(:C(H)Ph)(IPr)(PCy3) (9). A single-crystal x-ray diffraction study has been performed on 9. The thermal stability of 9 has been studied at 60° and its catalytic activity has been evaluated for the ring closing metathesis of di-Et diallylmalonate.

REFERENCE COUNT: 35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

...E + F ===> G...

L62 ANSWER 25 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

• c1-

G YIELD 47%

RX (2) RCT E 74663-75-5, F 50-00-0

STAGE(1) SOL 108-88-3 PhMe

STAGE (2)

RGT H 7647-01-0 HCl SOL 123-91-1 Dioxane

PRO G 250285-32-6 NTE PARAFORMALDEHYDE USED

RX(7) OF 15 COMPOSED OF RX(2), RX(3) RX(7) E + F ==> B

L62 ANSWER 25 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

B YIELD 79%

RX (3)

RX (2) RCT E 74663-75-5, F 50-00-0

STAGE(1) SOL 108-88-3 PhMe

STAGE(2) RGT H 7647-01-0 HCl SOL 123-91-1 Dioxane

PRO G 250285-32-6 NTE PARAFORMALDEHYDE USED

G 250285-32-6 K 865-47-4 t-BuOK B 244187-81-3 109-99-9 THF

L62 ANSWER 26 OF 35 CASREACT COPYRIGHT 2006 AGS on STN
ACCESSION NUMBER:
132:151738 CASREACT
TITLE:
Imidazolylidenes, imidazolinylidenes and imidazolidines
AUTHOR(S):
Acduengo, Anthony J., III; Krafczyk, Roland;
Schmutzler, Reinhard; Craig, Hugh A.; Goerlich, Jens
R.; Marshall, William J.; Unverzagt, Markus
Institut fur Anorganische und Analytiache Chemie, der
Techniachen Universitat - Carolo Wilhelmina,
Braunschweig, D-38106, Germany
SOURCE:
Tetrahedron (1999), 55(51), 14523-14534
CODDEN: TETRAB; ISSN: 0040-4020
PUBLISHER: Elsevier Science Ltd.
DOCUMENT TYPE: Journal
LANGUAGE: English
AB Starting from glyoxal and RNN12 (R = 2, 4, 6-Me3C6H2, 2, 6-(Me2CH)2C6H3), the
corresponding 1,3-diarylimidazolinium chlorides were obtained in a 3-step
sequence via dimimes and erbylenediamine dihydrochlorides. Subsequent
reduction with LIAIH4 furnished 1,3-diarylimidazolidines, while their
deprotonation with KH in TMF gave access to stable carbenes,
1,3-diarylimidazolin-2-ylidenes. Similarly substituted
imidazol-2-ylidenes are described for comparison.
REFFERNCE COUNT: 35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR
THIS

PORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE

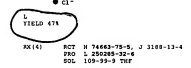
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

A + J ===> K... RX(3) OF 14

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

RCT A 56222-36-7, J 3188-13-4 PRO K 141556-45-8 SOL 109-99-9 THF RX (3)

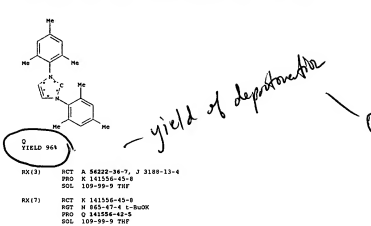
RX(4) OF 14 H + J ===> L... L62 ANSWER 26 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)



RX(9) OF 14 COMPOSED OF RX(3), RX(7) RX(9) A + J ===> Q

L62 ANSWER 26 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

L62 ANSWER 26 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)



RX(4) RCT H 74663-75-5, J 3188-13-4 PRO L 250285-32-6 SOL 109-99-9 THF

RX(5) RCT L 250285-32-6 RCT N 865-47-4 t-Buok PRO H 244187-81-3 SOL 109-99-9 THF

RX(10) OF 14 COMPOSED OF RX(4), RX(5)

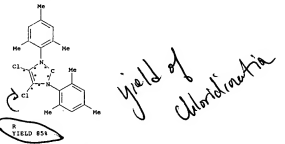
RX(10) H + J ===> H

i-Pr

RX(13) OF 14 COMPOSED OF RX(3), RX(7), RX(8) RX(13)  $A + J = P \times R$ 

L62 ANSWER 26 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

L62 ANSWER 26 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)



i-Pr Pr Cl N C i

RX(3) RCT A 56222-36-7, J 3188-13-4 PRO K 141556-45-8 SOL 109-99-9 THF

RX(7) RCT K 141556-45-9 RCT N 865-47-4 L-BuOK PRO Q 141556-42-5 SOL 109-99-9 THF

RX(8) RCT Q 141556-42-5 RCT P 56-23-5 CC14 PRO R 200730-48-9 SOL 109-99-9 THF

RX(14) OF 14 COMPOSED OF RX(4), RX(5), RX(6) RX(14) H + J ==> O

1-Pr N. 1-Pr 1-Pr C1 Pr-1

STEPS RX(4) RCT H 74663-75-5, J 3188-13-4
PRO L 250265-32-6
RCT L 250285-32-6
RCT N 865-47-4 t-BuoK
RX(5) RCT L 250285-32-6
RCT N 865-47-4 t-BuoK
RX(6) RCT M 244187-81-3
RCT M 244187-81-3
RCT P 56-23-5 CC14
PRO 0 258278-31-8
RCT N 56-23-9 THF

L62 ANSWER 27 OF 35 CASREACT COPYRIGHT 2006 ACS ON STN

ACCESSION NUMBER:

TITLE:

Efficient Cross-Coupling of Aryl Chlorides with Aryl Grignard Reagents (Kumada Reaction) Mediated by a Palladium/Inidazolium Chloride System

AUTHOR(S):

CORPORATE SOURCE:

Department of Chemistry, University of New Orleans, New Orleans, LA, 70148, USA
New Orleans, LA, 70148, USA
DOURLA of the American Chemical Society (1999), 121(42), 9889-9850

CODEN: JACSAT: ISSN: 0002-7863

American Chemical Society

Journal
LANGUAGE:

AB A general methodol. for the Kumada reaction was presented. In the presence of tris[µ-[(1,2-n;4.5-n]-(1E,4E)-1,5-diphenyl-1,4-pentadien-3-one]] dipalladium or palladium diacetate and a mindazolium chloride, aryl chlorides, aryl bromides or aryl iodides underwent a coupling reaction to give biphenyl derivs. Suitable imidazolium compds. Were 1,3-bis(2,4,6-trimethylphenyl)-1H-imidazolium chloride and 1,3-bis(2,6-bis(1-methylethyl)phenyl)-1H-imidazolium chloride and THERE ARE 64 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RX (2) OF 16 ...C + G ===> H

$$\begin{array}{c} 1-Pr \\ \\ \\ Pr-1 \\ \\ C \\ \end{array}$$

L62 ANSWER 27 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)



RX (2) RCT C 74663-75-5, G 50-00-0

STAGE(1) SOL 108-88-3 PhMe

STAGE (2) RGT I 7647-01-0 HC1 SOL 123-91-1 Dioxane

PRO H 250285-32-6 NTE paraformaldehyde used, prior prepns. were one-pot

L62 ANSWER 28 OF 35
ACCESSION NUMBER:
131:337140 CASREACT
ITILE:
131:337140 CASREACT
N,N'-Diferrocencylic Carbenes and Their
Derivatives
AUTHOR(S):
Bildstein, Benno: Malaun, Michael; Kopacka, Holger;
Wurst, Klaus: Mitterboeck, Martin; Ongania,

Opromolla, Giuliana: Zanello, Piero Institut fuer Allgemeine Anorganische und

CORPORATE SOURCE: Theoretische

Chemie, Universitaet Innsbruck, Innsbruck, A-6020,

Chemie, Universitaet Innsbruck, Innsbruck, A-6020,
Austria
Organometallics (1999), 18(21), 4325-4336
CODEN: ORGMD7: ISSN: 0276-7333
PUBLISHER: American Chemical Society
Journal
ANGUAGE: English
AB In continuation of the authors' work on Wanzlick/Arduengo carbenes
containing
redox-active ferrocenyl substituents the synthesis of N,N'-diferrocenyl
imidazol(in)ium salts as precursors of imidazol(in)-2-ylidenes is
reported. The necessary starting material for this chemical is
aminoferrocene, which was prepared by an improved and large-scale
synthesis
by the sequence solid lithioferrocene, iodoferrocene, Nferrocenylphthalimide, aminoferrocene. The preparation of
N,N'-diferrocenyl
heterocycles involves condensation of aminoferrocene with glyoxal to
afford N,N'-diferrocenyldiazabutadiene [Fc-DAB], reduction, condensation
with

of the formaldehyde, and oxidation with trityl salts to yield N,N'diferrocenylimidazol(in)ium salts. In situ deprotonation and trapping
with electrophiles yielded the expected metal complexes and derivs. in
some cases [Ag+ or S8], but attempted reaction with other transition
metals [e.g., Pd(II)] failed to give the corresponding complexes, due to
(i) steric hindrance by the two N-ferrocenyl substituents, (ii) reduced
acidity of the imidazol(in)ium precursors, and (iii) inaccessibility of
the free carbenes. Spectroscopic [IR, Raman, UV-visible, MS, NMR (IH,
13C, 109Ag)], structural [x-ray], and electrochem. [CV] properties are
reported and compared to those of other N-heterocyclic carbene derivs.

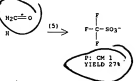
REFERENCE COUNT:

32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

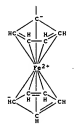
FORMAT

RX(5) OF 66 ...C + O + H ===> P... L62 ANSWER 28 OF 35 CASREACT COPYRIGHT 2006 ACS on STN



\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

PAGE 2-A



P: CM 2

PAGE 2-A

L62 ANSWER 28 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

L62 ANSWER 28 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

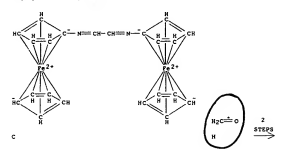
RX(5) RCT C 249644-26-6, O 1493-13-6

STAGE(1) RGT Q 557-20-0 Et2Zn SOL 75-05-8 MeCN, 110-54-3 Hexane

STAGE (2) RCT H 50-00-0

PRO P 249644-41-5 NTE PARAFORMALDEHYDE USED

RX(16) OF 66 COMPOSED OF RX(2), RX(3) RX(16) C + H ===> I



\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

HC H H CH

Fe 2+

TO H CH

TO H CH

Fe 2+

TO H CH

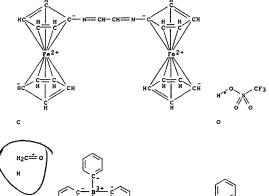
TO H

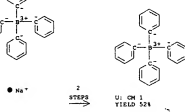
RX(2) RCT C 249544-26-6 RGT G 16853-85-3 LiAlH4 PRO F 249644-28-8

RX(3) RCT F 249644-28-8, H 50-00-0 PRO I 249644-30-2 SOL 67-64-1 Me2CO, 7732-18-5 Water

RX(18) OF 66 COMPOSED OF RX(5), RX(6) RX(18) C + O + H + T ===> t

L62 ANSWER 28 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

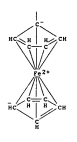




\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

L62 ANSWER 28 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Conti

PAGE 2-A



U: CM 2 YIELD 52%

RX(5) RCT C 249644-26-6, O 1493-13-6

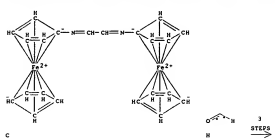
STAGE(1) RGT Q 557-20-0 Et22n SOL 75-05-8 MeCN, 110-54-3 Hexane

STAGE(2) RCT H 50-00-0

PRO P 249644-41-5 NTE PARAFORMALDEHYDE USED

RX(6) RCT P 249644-41-5, T 143-66-8 PRO U 249644-43-7 SOL 67-56-1 MeOH

RX(29) OF 66 COMPOSED OF RX(2), RX(3), RX(13) RX(29) C + H ===> AO L62 ANSWER 28 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

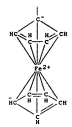


AO: CM 1

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

L62 ANSWER 28 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

PAGE 2-A



AO: CM 2 YIELD 78%

RX(2) RCT C 249644-26-6 RGT G 16853-85-3 LiAlH4 PRO F 249644-28-8

RX(3) RCT F 249644-28-8, H 50-00-0 PRO I 249644-30-2 SOL 67-64-1 Me2CO, 7732-18-5 Water

RX(13) RCT I 249644-30-2 RGT AP 341-02-6 Ph3C.BF4 PRO AO 249644-60-8 SOL 75-09-2 CH2C12

L62 ANSWER 29 OF 35 CASREACT COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER:
130:252064 CASREACT
TITLE:
A practical and efficient synthesis of
enantiomerically pure di-tert-butylethanediamine
Roland, Sylvain; Mangeney, Pierre; Alexakis, Alex
Lab. Chimie Organo-Elements, Univ. Pierre Marie
Curie,
SOURCE:
Synthesis (1999), (2), 228-230
CODEN: SYNTBF; ISSN: 0039-7881
Georg Thieme Verlag
DOCUMENT TYPE:
Journal
LANGUAGE:
English
AB A diastereoselective synthesis of 1,2-di-tert-butylethylenediamine was
developed by addition of Me3CMgCl to a chiral bis-imine derived from
glyoxal
and (S)-\alpha-methylbenzylamine. Addition of the bis-imine to the Grignard
reagent in hexane at 50° gave only one diastereomer detectable by
1H and 13C NMR. Hydrogenolysis of the phenylethyl groups led to the
expected free (R,R) diamine in good yields.
REFERENCE COUNT:
26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR
THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

PORMAT

RX(7) OF 10 COMPOSED OF RX(4), RX(2)
RX(7) 2 Q + C + H ===> I

RX(4) RCT Q 677-22-5

STAGE(1)
SOL 60-29-7 Et20, 110-54-3 Hexane

STAGE (2)
RCT C 138812-17-6
SOL 110-54-3 Hexane

STAGE (3)
RGT R 12125-02-9 NH4Cl
SOL 7732-18-5 Water

PRO G 221638-36-4
NTE stereoselective

RX(2) RCT G 221638-36-4, H 50-00-0

L62 ANSWER 29 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

STAGE(1)
 RGT E 64-18-6 HCO2H
 SOL 7732-18-5 Water

STAGE(2)
 RGT J 7732-18-5 Water
 SOL 60-29-7 Et20

STAGE(3)
 RGT K 584-08-7 K2CO3

PRO I 221638-37-5 NTE stereoselective (Continued)

L62 ANSWER 30 OF 35
ACCESSION NUMBER: 129:109035 CASREACT
TITLE: Stable tetraazafulvalenes. Synthesis and chemistry
AUTHOR(S): Kaepplinger, Christian: Beckert, Rainer: Imhof,
Wolfgang
CORPORATE SOURCE: Institut Organische Makromolekulare Chemie,
Friedrich-Schiller-Universitaet, Jena, D-07743,
Germany

Friedrich-Schiller-Universitaet, Jena, D-07743, Germany Journal fuer Praktische Chemie/Chemiker-Zeitung (1998), 340(4), 323-333 CODEN: JPCCEH; ISSN: 0941-1216 Johann Ambrosius Barth Journal

SOURCE:

PUBLISHER:

DOCUMENT TYPE: LANGUAGE: GI

The syntheses, properties and reactions of 1,3,6,7-tetrakis(arylamino)1,4,5,8-tetraarafulvalenes and their vinylogous derivs. are described.
The acylation of form- as well as acetamidine with bis-imidoyl chlorides
derived from oxalic acid formed reactive cyclic intermediates which
dimerized to tetraarafulvalenes I (X = (double bond) or bisvinylogous
tetraarafulvalenes I [X = (CH)2]. A further synthesis was found using
cycloacylation reaction of amidines with imidoyl chlorides followed by
prototropic migration of a-H. Thus, the vinylogous compound I (X =
(CH)4) and the phenylogous derivs. I (X = CHC6H4-2-CH, CHC6H4-4-CH) were
isolated in moderate to good yields. Besides amidines, other carboxylic
acid derivs. such as amides or thicamides were transformed into
corresponding tetraarafulvalenes. Due to the vicinal amino groups,
alkylation and acylation reactions were studied. For example, the
reaction with orthoformates yielded ring-fused products which may be
starting material for carbenes just as the cyclization product with
2. SCC12

Treatment of tetraazafulvalenes with anhydrous Fe(II) salts or Mo(CO)6 yielded deeply colored metal diazadiene complexes. Finally, reduction

metallic Li and subsequent alkylation constitutes a convenient synthetic entry to heterocyclic analogs of stilbene.

RX(8) OF 21 Q + 2 R ===> 8

L62 ANSWER 30 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

YIELD 45%

RCT RGT PRO Q 189115-08-0, R 75-11-6 D 121-44-8 Et3N S 210051-67-5 1330-20-7 Xylene RX (8)

L62 ANSWER 31 OF 35
ACCESSION NUMBER:
128:270277 CASREACT
COPYRIGHT 2006 ACS on STN
128:270277 CASREACT
Generation and trapping reactions of a formal 1:1
complex between singlet carbon and 2,2'-bipyridine
Weiss, Robert; Reichel, Silvia; Handke, Matthias;
Hampel, Frank
SOURCE:
SOURCE:
SOURCE:
PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
GI

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

AB Treatment of Ph3As+C(=N2)CO2CMe3 with CF3SO3H gave Ph3As+CH2OTf which with 2,2'-bipyridine (I) gave the diquat II which was in equilibrium with its conjugate base III (X-=OTf-); with excess I, II was converted to III

= OTf-). Ion exchange gave 75% III.H2O (X- = Br-) whose crystal structure

cure
was determined III (X- = Br-) in THF containing KOCMe3 and Se gave 100%

IV via the deprotonated singlet C compound V. The crystallog. extensive delocalization in IV, MO calcns. of III (X- = Br-) and V, isodesmic reactions and reactivity of V, and 13c and 1H NNR of these compds. are discussed. REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

RX(1) OF 5 A + B ==> C...

L62 ANSWER 31 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

• Br

VIELD 75%

RX (1) RCT A 205182-22-5, B 366-18-7

STAGE (1) SOL 75-05-8 MeCN

STAGE(2) SOL 60-29-7 Et20

STAGE (3)

RGT D 1643-19-2 Bu4N.Br SOL 75-09-2 CH2C12

PRO C 205182-29-2

RX(4) OF 5 COMPOSED OF RX(1), RX(2) RX(4) A + B + H ===> I

L62 ANSWER 31 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

RX (1) RCT A 205182-22-5, B 366-18-7

STAGE (1) SOL 75-05-8 MeCN

STAGE(2) SOL 60-29-7 Et20

STAGE (3)

RGT D 1643-19-2 Bu4N.Br SOL 75-09-2 CH2C12

PRO C 205182-29-2

RCT C 205182-29-2, H 1493-13-6 PRO I 205182-23-6 SOL 109-99-9 THF RX (2)

L62 ANSWER 32 OF 35 CASREACT COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 113:59021 CASREACT
Compounds with bridgehead nitrogen. Part 61.
Conformational equilibria in the

perhydrodipyrido[1,2-

AUTHOR (S):

CORPORATE SOURCE:

c:2',1'-e|imidazoles Banting, Lee: Crabb, Trevor A.: Fallah, Asadollah; Williams, Roger O. Dep. Chem., Portsmouth Polytech., Portsmouth/Hampshire, POl 2DT, UK Journal of Chemical Research, Synopses (1990), {1}, SOURCE:

CODEN: JRPSDC; ISSN: 0308-2342 Journal English

DOCUMENT TYPE: LANGUAGE:

syn-Perhydrodipyrido[1,2-c:2',1'-e]imidazole [I] has been shown to adopt an equilibrium in CDC13 solution at 25 °C between the enantiomeric N-outside-cis-syn-trans-conformers contrary to an earlier report [P. J., Chivers, et al., 1968) assigning a predominance of the trans-syn-trans-conformer. anti-Perhydrodipyrido[1,2-c:2',1'-e]imidazole [II] shows the expected preference for the trans-anti-trans-conformation.

RX(3) OF 3 COMPOSED OF RX(1), RX(2) RX(3) 2 A + 2 E ==> F + G

L62 ANSWER 32 OF 35 CASREACT COPYRIGHT 2006 ACS on STN

G YIELD 90% (50)

A 56100-22-2 C 1333-74-0 H2 B 23549-50-0 64-19-7 ACOH RX (1)

RX (2)

E 50-00-0, B 23549-50-0 F 22210-62-4, G 22210-68-0 7732-18-5 Water

L62 ANSWER 33 OF 35
ACCESSION NUMBER:
108:204028 CASREACT
Compounds with bridgehead nitrogen. 52. NMR spectra
and stereochemistry of the 2alkylperhydroimidazolo[3,4-a]pyridines
Banting, Lee; Crabb, Trevor A.
CORPORATE SOURCE:
50URCE:
50URCE:
696-706
CASREACT COPYRIGHT 2006 ACS on STN
108:204028 CASREACT
Compounds with bridgehead nitrogen. 52. NMR spectra
and stereochemistry of the 2alkylperhydroimidazolo[3,4-a]pyridines
Banting, Lee; Crabb, Trevor A.
Portsmouth Polytech.,
Pol SOURCE: 696-706

CODEN: MRCHEG; ISSN: 0749-1581 Journal English

DOCUMENT TYPE: LANGUAGE: GI

AB In contrast to perhydrooxazolo[3,4-a]pyridine and perhydrothiazolo[3,4-a]pyridine, which adopt equilibrium in CDC13 solution at room temperature containing ca 70% trans fused conformers in equilibrium with O- or S-inside cis fused conformers,

2-alkylperhydroimidazolo[3,4-a]pyridines I [R = Rl = H, R2 = Me, Pr, (CR2)5Me, (CR2)3CHMe(CR2)3CHMe2, CHMe2, cyclohexyl, CMe3; R = Et, Rl = H, R2 = Me, cyclohexyl, CMe3; R = H, R1 = Me, R2 = Me (CR2)5Me, (CMC)3CHMe2, CMC)3CMC are found to adopt equilibrium containing >98% trans fused conformers.

comparison of NMR parameters of I (R = Rl = H, R2 = Me) with those of the 2 isomers of I (R = H, R1 = R2 = Me) indicates an equilibrium for the former compound

compound

between the two trans fused conformers, with ca 83% of that conformation

containing a trans arrangement of nitrogen lone pairs. These

observations are

explained in terms of the generalized anomeric effect.

RX(54) OF 62 COMPOSED OF RX(14), RX(28), RX(11) RX(54) AI ===> AA + AB

T. (10)

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L62 ANSWER 33 OF 35 CASREACT COPYRIGHT 2006 ACS ON STN RX(14) RCT AI 16273-56-6 RCT AK 16940-66-2 NaBH4 PRO AJ 114366-07-3 SOL 67-56-1 MeOH
                                                                                                                                                        (Continued)
                                   AJ 114366-07-3

AR 1333-74-0 H2

Z 114366-21-1

1314-15-4 PtO2

64-19-7 ACOH
                         RCT
RGT
PRO
CAT
RX (28)
                        RCT Z 114366-21-1
RGT C 50-00-0 HCHO
PRO AA 114365-95-6, AB 114365-98-9
SOL 7732-18-5 Water
RX (11)
RX(56) OF 62 COMPOSED OF RX(15), RX(29), RX(12) RX(56) AM \Rightarrow\Rightarrow AD + AE
                                                                                                                               (CH<sub>2</sub>)5
                                                             STEPS
                                      (CH2)5
ΑE
                                   AM 114366-24-4

AK 16940-66-2 NaBH4

AN 114366-08-4

67-56-1 MeOH
RX (15)
                                   AN 114366-08-4
AR 1333-74-0 H2
AC 114366-22-2
1314-15-4 PtO2
64-19-7 ACOH
                        RCT
RGT
PRO
CAT
SOL
RX (29)
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L62 ANSWER 33 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

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RX(12) RCT AC 114366-22-2

RX(58) OF 62 COMPOSED OF RX(16), RX(30), RX(13)

RX(58) AO ===> AG + AH

RX(16) RCT AC 107954-71-2
RCT AC 107954-71-2
RCT AC 107954-71-2
RCT AC 16940-66-2 NaBH4
PRO AP 114366-09-5
SOL 67-56-1 MeOH

RX(30) RCT AP 114366-09-5
RGT AR 1333-74-0 H2
PRO AP 114366-09-5
RGT AR 1134-15-4 PtO2
SOL 64-19-7 ACOH

RX(13) RCT AF 114366-23-3
RGT C 50-00-0 HCHO
RX(13) RCT AF 114366-09-5
RGT AR 1314-15-4 PtO2
SOL 64-19-7 ACOH

RX(13) RCT AF 114366-23-3
RGT C 50-00-0 HCHO
RX(13) RCT AF 114366-20-5
RGT C 50-00-0 HCHO
RX(13) RCT AF 114366-23-3
RGT C 50-00-0 HCHO
RX(13) RCT AF 114366-29-5
RGT C 50-00-0 HCHO
RX(13) RCT AF 114366-23-3
RGT C 50-00-0 HCHO
RX(13) RCT AF 114366-23-3
RGT C 50-00-0 HCHO
RX(13) RCT AF 114366-23-3
RGT C 50-00-0 HCHO
RX(13) RCT AF 114366-97-8, AH 114366-00-6
SOL 7732-18-5 Water
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L62 ANSWER 34 OF 35
ACCESSION NUMBER:
TITLE:
Synthesis of 1,3-disubstituted diazolidines
Lambert, Joseph B.; Huseland, Dave E.; Wang, Gen Tai
Dep. Chem., Northwestern Univ., Evanston, IL, 60201,
USA
SOURCE:
Synthesis (1986), (8), 657-8
CODEN: SYNTBF; ISSN: 0039-7881
Journal
LANGUAGE:
GI

RN NR1

AB Sym. and unsym. RNHCH2CH2NHR1 (I; R = PhCH2, Et, Ph; Rl = Me, Ph, CH2Ph) were obtained by the reduction of RNHCOCONHR1 (II) with LiAlH4. II were readily produced by treatment of di-Et oxalate with primary amines. I gave imidazolidines III on treatment with CH2O.

RX(11) OF 14 COMPOSED OF RX(3), RX(4) RX(11)  $\mathbf{F}$  + K ===>  $\mathbf{L}$ 

RX(3) RCT F 7666-51-5 RGT I 16853-85-3 LiAlH4 PRO H 56904-09-7 SOL 109-99-9 THF

RX(4) RCT K 50-00-0, H 56904-09-7 PRO L 105900-08-1 SOL 64-17-5 EtOH, 7732-18-5 Water

L62 ANSWER 35 OF 35
ACCESSION NUMBER:
TITLE:
Models for tetrahydrofolic acid. I. Condensation of formaldehyde with tetrahydroquinoxaline analogs
Benkovic, Stephen J.: Benkovic, Patricia A.: Comfort, David R.
CORPORATE SOURCE:
SOURCE:
Dovid R.
Pennsylvania State Univ., University Park, PA, USA Journal of the American Chemical Society (1969), 91(19), 5270-9
CODEN: JACSAT: ISSN: 0002-7863
Journal and State Univ. Benkovic, Patricia A.: Comfort, David R.
Pennsylvania State Univ., University Park, PA, USA JOURNAT TYPE:
LANGUAGE:
English
AB To investigate the mechanisms of tetrahydrofolic acid catalyzed one carbon unit transfers, we have synthesized several tetrahydroquinoxaline

unit transfers, we have synthesized several tetrahydroquinoxaline analogs.

A kinetic investigation of the condensation with CH2O of one of these models reveals the intermediacy of the iminium cation as a steady-state species and the importance of general catalysis in formation of the imidazolidine ring, the latter a model for 5,10-methylene tetrahydrofolic acid. The relevance of these results to the mechanism of one carbon unit transfers and the importance of certain structural and electronic features
in the actual cofactor is discussed.

RX(4) OF 4 COMPOSED OF RX(1), RX(3) RX(4) A + E ===> P

L62 ANSWER 35 OF 35 CASREACT COPYRIGHT 2006 ACS on STN (Continued)

F YIELD 70%

RX (1)

RCT A 62294-77-3
RGT C 16940-66-2 NaBH4
PRO B 23792-11-2
SOL 111-96-6 (MeOCHZCH2)20
NTE Classification: Chemoselective; Dearomatisation; Reduction; \$
Conditions: NaBH4; diglyme 1h; 20 deg

RX (3)

B 23792-11-2, E 50-00-0 F 25187-69-3 123-91-1 Dioxane, 7732-18-5 Water Classification: Heterocycle formation; Condensation; N-Alkylation; # Conditions: 1,4-dioxan H2O; heat water bath